# Sensitivity Analysis of a Mixed-Phase Chemical Mechanism using Automatic Differentiation

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#### **Abstract**

A mixed-phase chemistry box model is applied to study heterogeneous chemistry and its effect on tropospheric gas-phase chemistry, particularly on photochemical production of  $O_3$  and photochemical indicators for  $O_3$ -NO<sub>x</sub>-hydrocarbon sensitivity, under a variety of atmospheric conditions ranging from remote marine to heavily-polluted atmospheres. A subsequent sensitivity analysis of the mixed-phase chemical mechanism is conducted using the novel automatic differentiation ADIFOR tool, which calculates the local sensitivities of species concentrations in gas, aqueous and aerosol phases with respect to a variety of model parameters. The main chemical reaction pathways in all phases, interfacial mass transfer processes, and ambient physical parameters that affect tropospheric  $O_3$  formation,  $O_3$ -precursor relations and photochemical indicators under all modeled conditions are identified and analyzed.

The results show that the presence of clouds and aerosols not only reduces many gas-phase species concentrations and the total oxidizing capacity but alters O<sub>3</sub>-precursor relations. Cloud chemistry is the dominant heterogeneous process under the remote and marine atmospheres. Aerosols are important scavengers for gaseous species in polluted atmospheres when the total aerosol surface area is larger than 1000 µm<sup>2</sup> cm<sup>-3</sup>. Decreases in concentrations and formation rates of O<sub>3</sub> can be up to 27% and 100%, respectively, depending on the initial atmospheric conditions and preexisting surfaces. The significant decreases in photochemical O<sub>3</sub> formation are primarily caused by the aqueous-phase reactions of O<sub>2</sub>- with dissolved HO<sub>2</sub> and O<sub>3</sub> under most cloudy conditions and heterogeneous uptake of O<sub>3</sub>, HCHO, HO<sub>2</sub>, H<sub>2</sub>O<sub>2</sub> and NO<sub>2</sub> in the presence of aerosols. Heterogeneous chemical processes also affect the photochemical indicators and their sensitivities to many model parameters in a variety of ways.

#### 1. Introduction

Air pollution models have provided the most effective means to quantify the impact of human activities on the atmosphere and to develop balanced and cost-effective emission control strategies for air pollutants such as tropospheric ozone (O<sub>3</sub>) during the last several decades. The effectiveness of these abatement strategies depends on the reliability of model predictions, which is further built upon the accurate understanding and appropriate parameterization of a variety of atmospheric processes in current air quality models. Sensitivity analysis of these models is a valuable diagnostic tool capable of identifying the most influential model parameters and predicting the changes in the responses due to arbitrary perturbations in the parameters. In particular, it can evaluate the relative importance of various physical and chemical processes, the influence of the model input and mechanistic parameters, and the role of individual species and their correlations in the atmospheric system as a whole, thus providing scientific insights and suggesting research priorities for laboratory and field studies. While sensitivity analysis of 3-D models can acquire scientific information that is useful in long range scientific and regulatory analysis regarding air pollutant abatement, sensitivity analysis of atmospheric chemistry box models provides scientific insights into the detailed chemical mechanisms, on which the condensed mechanisms typically used for 3-D models are based.

Among many uncertain model mechanistic and input parameters, the representation of atmospheric chemistry used in the models is of large uncertainty. While the basic set of reactions that leads to O<sub>3</sub> production has been identified, different chemical representations produce different O<sub>3</sub> predictions. For a given chemical mechanism containing a large number of chemical reactions, model predictions of key species are sensitive to the rate parameters and product yields, which may have large uncertainties when they are based upon limited experiments or estimation (Stockwell et al., 1993). The uncertainties of the system will greatly increase in the presence of various atmospheric condensed phases (e.g., clouds, aerosols, fog, rain, and snow). A growing number of studies have shown the importance of these heterogeneous processes. For example, the heterogeneous oxidation of SO<sub>2</sub> on aerosols may account for nearly 60% of the tropospheric sulfate formation in marine environments (Luria and Sievering, 1991). Aerosols may be an important sink for HO<sub>2</sub> and influence the tropospheric H<sub>2</sub>O<sub>2</sub> level (Ross and Noone, 1991). Clouds may decrease the gas-phase NO<sub>x</sub>, O<sub>3</sub>, OH, HO<sub>2</sub> and HCHO by 10-85 % in the global troposphere (Lelieveld and Crutzen, 1990). The decrease in O<sub>3</sub> formation is due to aqueous-phase radical reactions in typical urban polluted atmospheres (Walcek et al., 1997). However, many factors such as the reaction probability and product yields for these important heterogeneous reactions remain unknown, which creates several orders of magnitude higher uncertainties in mixed-phase chemistry than in gas-phase chemistry.

Rigorous sensitivity and uncertainty analyses of chemical mechanisms in 3-D comprehensive chemistry and transport models are rarely performed because of the computational limitation. Previous studies on chemical mechanisms and their sensitivity using 0-2D models have largely focused on limited numbers of species and model parameters for a given chemical mechanism that contains either gas-phase or aqueous-phase chemistry only. Milford et al. (1992) compared two gas-phase mechanisms: the LCC of Lurmann et al. (1987) and the CBM-IV of Gery et al. (1989) through evaluating sensitivities of model-predicted O<sub>3</sub>, H<sub>2</sub>O<sub>2</sub> and HCHO to initial concentrations and various reaction rate constants. Gao et al.

(1995) conducted a sensitivity and uncertainty analysis for the gas-phase mechanism in the second-generation Regional Acid Deposition Model (RADM2, Stockwell et al., 1990) and identified the most influential reactions to overall model predictions. Pandis and Seinfeld (1989) developed a full aqueous-phase mechanism for cloud chemistry and derived a condensed mechanism through sensitivity analysis without simulating gas-phase reactions. Since there are few models capable of realistically simulating atmospheric heterogeneous processes due to many uncertain parameters and pathways associated with these processes, the sensitivity of overall model predictions to these heterogeneous processes has been seldom quantified. To improve our understanding of heterogeneous chemistry and the associated uncertainties, a state of the art representation of all known or suspected heterogeneous chemical processes needs to be incorporated into existing atmospheric chemical mechanisms. A comprehensive sensitivity analysis based on such a mixed-phase chemical mechanism will be extremely valuable for investigating the effect of heterogeneous chemistry on photochemical processes and providing guidance for field and laboratory studies of important heterogeneous chemical reactions.

In this paper, we applied a mixed-phase chemistry box model MaTChM (Zaveri, 1997; Zhang et al., 1996) to study the effects of heterogeneous processes associated with clouds and aerosols on tropospheric gas-phase chemistry. A subsequent comprehensive sensitivity study of MaTChM was conducted using the automatic differentiation ADIFOR 2.0 tool (Bischof et al., 1992, 1996a). The sensitivities of individual species and photochemical indicators with respect to a variety of model input and mechanistic parameters are calculated and evaluated under a broad range of atmospheric conditions. Our primary objectives are to (i) quantify the impact of clouds and aerosols on tropospheric photochemistry; (ii) evaluate the sensitivity of overall model predictions to model parameterizations and inputs in the absence/presence of clouds and aerosols; (iii) identify the most important reactions in the condensed phases and the most influential model parameters under typical atmospheric conditions. The particular emphasis will be directed towards the evaluation of chemical pathways and parameters that affect tropospheric O<sub>3</sub> formation, O<sub>3</sub>-precursor relations and the measurable "indicators" that indicate the sensitivity of O<sub>3</sub> to emissions of nitrogen oxides (NO<sub>x</sub>) or reactive organic gases (ROG).

## 2. Mixed-Phase Chemistry Box Model

#### 2.1 Mixed-Phase Chemical Mechanism

The time-dependent mixed-phase <u>Mass Transfer</u> with <u>Chemical Reactions Box Model (MaTChM)</u> treats gas and aqueous phase chemistry, heterogeneous surface reactions on aerosols, and the dynamic exchange of species between gas and condensed phases in a homogeneous parcel of air containing cloud droplets and aerosol particles. Emission, dilution, transport and deposition processes are neglected. The model includes 57, 58, and 29 species in the gas, aqueous and aerosol phases, respectively, as shown in Table 1. The mixed-phase chemical mechanism is designed to be applicable in a variety of atmospheric conditions ranging from remote marine to heavily-polluted. It consists of 125 gas-phase reactions, 120 aqueous-phase reactions, 47 dissolution and dissociation equilibria and 29 aerosol surface reactions. The gas and aqueous-

phase chemistry was taken from the earlier version of MaTChM developed by Zaveri (1997), in which a condensed dimethyl sulfide (DMS) chemistry was derived based on the comprehensive DMS chemistry of Yin et al. (1990) and combined with the condensed Carbon Bond Mechanism IV(CBM-IV) of Gery et al. (1989) and the full aqueous-phase chemistry of Pandis and Seinfeld (1989). In Zaveri (1997), the combined gas and aqueous phase chemistry was further modified for simulations in relatively clean atmospheres through explicitly treating the lesser reactive methane and ethane, and their intermediate radicals in CBM-IV and adding the aqueous-phase oxidations of DMS and DMSO by dissolved oxidants. In addition, the mass transfer between gas and aqueous phases was simulated based on the two-resistance theory of Astarita (1967). A list of gas and aqueous-phase reactions taken from the earlier work of Zaveri (1997) is given in Appendix A-1 through A-4. It is noted that these mechanisms are partially different from the current work of Zaveri (1997) in which some rate constants have been updated and additional reactions have been included. In this work, we extended the earlier version of MaTChM of Zaveri (1997) to include heterogeneous uptake of gaseous species on aerosols to explore the effect of aerosols on photochemistry and to compare it with the effect of clouds. The heterogeneous uptake on aerosols is treated as a pseudo first-order mass transfer process (Li et al., 1993; Dentener et al., 1996). The model parameterization is described in detail below.

## 2.2 Mathematical Description

The governing equation for the mixed-phase chemical system can be given as:

$$\frac{d\overline{G}_i}{dt} = P_{Gi} - D_{Gi}\overline{G}_i - H_iK_{Li}(\frac{3}{a})V_c(\overline{G}_i - \frac{\overline{A}_iN_{avo}}{1000H_i}) - K_{pi}\overline{G}_i$$
(1)

$$\chi \frac{d\overline{A_i}}{dt} = \chi(P_{Ai} - D_{Ai}\overline{A_i}) + H_i K_{Li} \left(\frac{3}{a}\right) V_c \left(\overline{G_i} - \frac{\overline{A_i} N_{avo}}{1000 H_i}\right)$$
 (2)

$$Y\frac{d\overline{P_i}}{dt} = K_{pi}\overline{G_i} \tag{3}$$

where,  $\overline{G}_i$  is the bulk or average gas-concentrations of species i,  $\overline{A}_i$  and  $\overline{P}_i$  are the average aqueous- and aerosol-phase concentrations of species i, respectively (defined as the total amount of species in the aqueous or aerosol phases divided by the total volume of water in clouds or aerosols, respectively).  $P_{Gi}$ ,  $D_{Gi}$ ,  $P_{Ai}$ , and  $D_{Ai}$  are the chemical production and destruction terms for the gas and aqueous phases, respectively. a is the radius of cloud droplets which is assumed to be monodisperse,  $H_i$  is the effective Henry's Law constant,  $V_c$  is the liquid water content, and  $N_{avo}$  is Avogadro's number.  $K_{Li}$  is the overall mass transfer coefficient accounting for gas-phase diffusion, interfacial mass transfer, aqueous-phase diffusion and reactions in clouds;  $K_{pi}$  is the pseudo-first-order rate coefficient which describes the overall net heterogeneous loss rate of gas-phase species on aerosol surfaces.  $\chi$  is the conversion factor from moles L-1 to molecules cm<sup>-3</sup>, =10<sup>-3</sup>  $N_{avo} V_c$ . Y is the conversion factor from  $\mu g$  m<sup>-3</sup> to molecules cm<sup>-3</sup>, = 10<sup>-9</sup> ( $R T C_{air}/P M_i$ ), where R is the gas

constant in atm m<sup>3</sup> (Kmol K)<sup>-1</sup>, T is the temperature in K, P is the pressure in atm,  $C_{air}$  is the air concentration in molecules cm<sup>-3</sup>, and  $M_i$  is the molecular weight of species i.

 $K_{Li}$  can be calculated based on the two-film theory of Astarita (1967) (Zaveri, 1997):

$$\frac{1}{K_{Li}} = \frac{1}{k_{li}} + \frac{H_i}{k_{gi}} \tag{4}$$

with 
$$\frac{1}{k_{gi}} = \frac{a}{D_{gi}} + \frac{4}{\alpha_i \overline{\nu}}$$
 and  $k_{li} = \frac{D_{li}}{\delta_{li}}$ 

where  $k_{li}$  is the liquid-side mass-transfer coefficient with chemical reaction for species i;  $k_{gi}$  is the combined gas-phase and interfacial mass-transfer coefficient for species i;  $D_{gi}$  and  $D_{li}$  are the gas-phase and liquid-phase diffusion coefficients for species i, respectively;  $\alpha_i$  is the mass accommodation coefficient (i.e., sticking coefficient) for species i; v is the mean molecular speed; and  $\delta_{li}$  is the liquid-side hypothetical film thickness for species i. It depends on the cloud droplet radius and the effective aqueous-phase first-order reaction rate constant,  $D_{Ai} = P_{Ai}/\overline{A_i}$ . A detailed description of the mass transfer treatment can be found in Zaveri (1997).

The net removal rate on aerosols,  $K_{pi}$ , can be given as (Heikes and Thompson, 1983):

$$K_{pi} = \int_{r_1}^{r_2} k_{di}(r) \cdot n(r) dr \tag{5}$$

where  $k_{di}(r)$  is the gas-to-particle diffusion rate constant of species i for a particle of radius r. It can be calculated by the Fuchs and Sutugin interpolation equation (Fuchs and Sutugin, 1970):

$$k_{di} = \frac{4\pi r D_{gi} V}{1 + K_n (\lambda + 4(1 - \gamma_i) / 3\gamma_i)}$$
(6)

where the variable V represents ventilation factor which is close to 1,  $K_n$  is the Knudson number, defined as the ratio of the effective mean free path of a gas molecule in air,  $\lambda$ , to the particle radius r,  $\gamma_i$  is the uptake coefficient (i.e., reaction probability) for species i. n(r)dr is the aerosol number density between radii r and r+dr. It can be described using the lognormal size distributions for three aerosol modes (i.e., nucleation  $(0.001\text{-}0.1 \ \mu\text{m})$ , accumulation  $(0.1\text{-}1 \ \mu\text{m})$ , and coarse modes  $(>1 \ \mu\text{m})$ ) (Jaenicke, 1993):

$$\frac{dN(r)}{d(\log r)} = \sum_{j=1}^{3} \frac{n_j}{\sqrt{2\pi} \log \sigma_j} \exp\{\frac{-(\log r/R_j)^2}{2(\log \sigma_j)^2}\}$$
 (7)

where N(r) is the cumulative number concentration for particles with radius larger than r,  $n_j$ ,  $R_j$  and  $\sigma_j$  are the number density, mean radius and standard deviation of mode j, respectively.

#### 2.3 Mass Accommodation Coefficient and Uptake Coefficient

The mass accommodation coefficient  $\alpha$  is a fundamental parameter for heterogeneous uptake of gaseous species on particles. It represents the probability that a gas molecule reaching the particle surface will adhere to it. The sticking molecule may remain on or penetrate the surface or be desorbed back to the gas phase. The net flux to the surface can be described by the uptake coefficient  $\gamma$  (i.e., reaction probability), which is defined as the fraction of collisions that leads to irreversible loss of gaseous species on the particle surface. The distinction between  $\alpha$  and  $\gamma$  becomes important when desorption occurs.  $\gamma$  represents a lower limit to  $\alpha$  when the various processes can not be separated. Although  $\alpha$  and  $\gamma$  for various species have been extensively measured during the last several years, there are still large discrepancies among measured values, with the uncertainty factors ranging from 2-3 and 2-10, respectively (DeMore et al., 1994). Thus, a constant value is usually assumed for all species in current model studies (e.g., Pandis and Seinfeld, 1989; Dentener, 1993; Li et al., 1993). Since the magnitudes of  $\alpha$  or  $\gamma$  for various species vary from 10-8 to 1, however, this simplified treatment may introduce significant errors when estimating the effects of heterogeneous processes on photochemistry.

In this study, we incorporate the best-guessed individual mass accommodation coefficients,  $\alpha_i$ , either from measurements or indirectly derived from kinetic measurements, to calculate the overall mass transfer coefficients  $K_{Li}$  between gas-phase and cloud droplets, as shown in Eq. (4). In the light of the increasing measurements of  $\alpha_i$  and  $\gamma_i$ , it is assumed that all gaseous species that can be transferred to cloud droplets may undergo similar heterogeneous uptake and reactions on deliquescent aerosols. As an initial study indicating the effect of aerosols on tropospheric chemistry, these removal processes are assumed to be irreversible. The individual uptake coefficients  $\gamma_i$  measured or assumed based on the measured  $\alpha_i$  are used to calculate the irreversible heterogeneous loss rate  $K_{pi}$  of gases on aerosols, as shown in Eq. (5). Table 2 and 3 show  $\alpha_i$  and  $\gamma_i$  of condensing species used in the model and the resulting  $K_{pi}$  under the five modeled conditions, which will be described in the next section.  $K_{pi}$  is a strong function of available aerosol surfaces. It ranges from 1.0 x  $10^{-2}$  to  $1.0 \times 10^{-10}$  s<sup>-1</sup> and is up to 2-3 orders of magnitude higher under the polluted conditions than those under relatively clean conditions.

## 2.4 Model Inputs and Simulation Approaches

The sensitivities of overall model predictions are calculated along with the base simulations under a broad range of atmospheric conditions including heavily-polluted (Heavily-polluted), polluted (Urban), moderately-polluted (Rural), marine (Marine) and remote marine (Remote) conditions. The initial conditions are given in Table 4. Most initial concentrations are mainly based on literature values and the initial composition of hydrocarbon species is from Stockwell et al. (1990). The initial NO<sub>x</sub> concentrations range from 0.1 ppb under the Remote conditions to 110 ppb under the Heavily-polluted conditions, and the corresponding initial ROG ranges from 1.0 to 1100 ppb, with a ratio of ROG/NO<sub>x</sub> of 30 for the Urban conditions and 10 for other conditions. Under each condition, we further evaluate four scenarios: clear air (Gas) with only gas-phase reactions included, gas+aerosol (Aerosol) with gas-phase and aerosol surface

reactions included, gas+cloud (Cloud) with gas- and aqueous-phase reactions included, and gas+cloud+aerosol (Cloud+Aerosol) with all reactions included. A constant cloud droplet radius of 10 μm and a volumetric water content of 4 x 10<sup>-7</sup> are assumed for all simulations under cloudy conditions. These values are based on the measurements of the most common cloud types (e.g., stratus and cumulus) in the boundary layer and the free troposphere (Pruppacher and Klett, 1980; Heymsfield, 1993). The resulting total droplet number density is about 95 cm<sup>-3</sup>, with total surface area of 1.2 x 10<sup>-5</sup> μm<sup>2</sup> cm<sup>-3</sup>. Different aerosol size distributions measured under different atmospheric conditions (e.g., Whitby, 1978, Jaenicke, 1993 and Fitzerald, 1991) are used in this work. The total aerosol surface area available for surface uptake varies from a few tens μm<sup>2</sup> cm<sup>-3</sup> under relatively clean conditions to a few thousands μm<sup>2</sup> cm<sup>-3</sup> in polluted atmospheres.

All simulations began at local noon and were run for 2 hours. During this period of time, the atmospheric mixing, generally in equilibrium with emissions, has minimum effect on distribution and ambient concentrations of species. With maximum photolysis rates, peak ozone levels can be reached at approximate 2-4 p.m. From gas-aqueous mass transfer standpoint, the gas and aqueous-phase equilibria can be established as rapid as within a few minutes and a 2-hr period is sufficient to simulate the aqueous-phase reactions and predict their feedback to gas-phase chemistry, as shown in Pandis and Seinfeld (1989). A constant temperature of 280 K and a relative humidity (R.H.) of 80% are used for all simulations. While the high R.H. is associated with most cloudy conditions and pollution episodes, it also warrants ambient aerosols in deliquescent state with certain amounts of liquid water on the surface. A temperature of 280 K reflects a lower limit of the observed values in the boundary layer clouds. The photolysis rates are calculated continuously throughout the simulation period at a latitude of 40° N under summer time conditions. Clouds and aerosols are assumed to remain all the time for Aerosol, Cloud and Cloud+Aerosol cases. The set of ordinary differential equations (ODEs) describing the mass balances is solved using LSODE (Livermore Solver for Ordinary Differential Equations) software package (Hindmarsh, 1983). An integration time step of 1 minute is used.

#### 3. Sensitivity Analysis Method

The system of ODEs for gas, aqueous and aerosol phase concentrations shown in Eq. (1) through (3) can be written in a generalized form:

$$\frac{dc_k}{dt} = r_k(\mathbf{c}, t; \mathbf{q}) \quad \mathbf{c}(t_0) = \mathbf{c}_0$$
 (8)

where  $\mathbf{c}$  is the N-dimensional vector of concentrations,  $\mathbf{q}$  is the M-dimensional time-independent vector of model parameters,  $c_k$  is the concentration of the  $k^{th}$  species,  $r_k$  is the reaction rate of the  $k^{th}$  species and is dependent on  $\mathbf{c}$ ,  $\mathbf{q}$ , and the time t,  $\mathbf{c}_0$  is the initial condition on  $\mathbf{c}$ .

The sensitivities of the concentration of the  $k^{th}$  species with respect to the  $l^{th}$  parameter

$$S_{kl} = \frac{\partial c_k}{\partial q_l} \tag{9}$$

can be computed using a variety of techniques including the brute force method, Green's function analysis (Cho et al., 1987), adjoint Green's function analysis (Hall and Cacuci, 1983), and several variations of the direct decoupled methods (Dunker, 1984). Most sensitivity studies of 3-D atmospheric models have been conducted using the so-called brute force method, i.e., a number of model parameters are varied and the simulation results are then compared. This approach is closely related to finite-difference approximations of derivatives, where the derivatives  $\partial c_k/\partial q_l$  are approximated as  $[c_k(q+he_l)-c_k(q)]/h$ , where  $e_l$  is a vector that is all zero except for an entry of unity in its  $l^{th}$  component. This method becomes less viable as the model becomes more comprehensive because of the large computational burden. In addition, a suitable choice of the stepsize h may not be obvious. An alternative technique for sensitivity study is automatic differentiation (AD). This approach is superior to finite difference approximation of the derivatives because the numerical values of the computed derivatives are guaranteed to be accurate to machine precision and the computational effort can be significantly lower (Griewank and Corliss, 1991). In our work, we employ the ADIFOR (Automatic DIfferentiation in FORTRAN) tool (Bischof et al., 1992, 1996a), to compute a complete set of sensitivities for all species with respect to all model parameters.

#### 3.1 Introduction for ADIFOR

Automatic differentiation is an approach to computing sensitivities which relies on the fact that every function, no matter how complicated, is executed on a computer as a sequence of elementary operations such as additions, multiplications, and elementary functions such as *sin* and *cos* (e.g., Rall, 1981; Griewank and Corliss, 1991; Berz et al., 1996). By applying the chain rule of derivative calculus over and over again to the composition of those elementary operations, one can compute, in a completely mechanical fashion, derivatives of a function *f* that are correct up to machine precision. Traditionally, two approaches to AD have been developed: the so-called forward and reverse modes, which are distinguished by how the chain rule is used to propagate derivatives through the computation. The two approaches are briefly summarized below. A detailed description can be found in Bischof et al. (1996a) and the references therein.

Assume that we have a code for the computation of a function f,  $f:q \in R^M \mapsto c \in R^N$  we wish to compute  $S_u$  in Eq. (9), i.e., the derivatives of the dependent variables  $c_k$  with respect to independent variables  $q_l$ . The forward mode propagates derivatives of intermediate variables with respect to the independent variables and follows the control flow of the original program. By exploiting the linearity of differentiation, the forward mode allows us to compute arbitrary linear combinations  $J \cdot U$  of columns of the Jacobian J = dc / dq. For an  $N \times p$  matrix U, the effort required is roughly O(p) times the runtime and memory of the original program. In particular, when U is a vector u, we compute the directional derivative  $J * u = \lim_{k \to 0} \frac{f(q + h \cdot u) - f(q)}{h}$ .

In contrast, the reverse mode of AD propagates derivatives of the final result with respect to an intermediate quantity, in a fashion that is closely related to discrete adjoint techniques. To propagate adjoints, one must be able to reverse the flow of the program, and remember or recompute any intermediate value that nonlinearly affects the final result. However, if these technical difficulties can be overcome, for a

 $z \times M$  matrix W, the reverse mode allows us to compute the row linear combination  $W \cdot J$  with O(z) times as many floating-point operations as required for the evaluation of f. When W is a row vector w, we compute the derivative  $\partial (w^T * c)/\partial \mathbf{q}$ .

The ADIFOR 2.0 system, jointly developed by Argonne National Laboratory and Rice University (Bischof et al., 1996a), provides the directional derivative computation possibilities associated with the forward mode of AD. ADIFOR has several novel features:

- It employs a hybrid scheme that combines the forward mode overall with the reverse mode at the statement level;
- The SparsLinC subsystem allows transparent exploitation of sparsity in derivative objects, without any apriori knowledge of sparsity structure (Bischof et al., 1996b);
- The ADIntrinsics subsystem provides complete and user-configurable handling of all Fortran 77 intrinsics;
- ADIFOR supports almost all of Fortran 77, in particular, the differentiation of codes using any or all of the real or complex single or double precision-valued data types.

ADIFOR has been applied to sensitivity analysis in a variety of problem domains as described in Bischof et al. (1996a). In particular, ADIFOR has been used for sensitivity analysis of atmospheric models including 0-D Sulfur Transport Eulerian Model (STEM-II) (Carmichael et al., 1997) and MM5, a 3-D finite-difference weather model for meso- and regional-scale studies (Bischof et al., 1996c). Application of ADIFOR to a code requires that the user designate the independent and dependent variables with respect to differentiation, and provide the location of the source files containing the subroutines to be differentiated. ADIFOR then generates a new FORTRAN code that computes both the original simulation and its derivatives. The user then provides a derivative driver that selects at runtime what derivative is to be computed (this corresponds to the instantiation of the matrix *U* mentioned above) and invokes the ADIFOR-generated code. Similar functionality has been recently developed for ANSI-C programs with the ADIC tool (Bischof et al., 1996d) and that a survey of available automatic differentiation tools can be found at http://www.mcs.anl.gov/Projects/autodiff/AD\_Tools.

## 3.2 Application of ADIFOR to MaTChM

To compute the sensitivities  $S_{kl}$  in Eq. (9), we applied ADIFOR directly to the MaTChM code, which basically consists of the code for LSODE and the code for rate equations. That is, we considered the MaTChM code as a "black box" that, given an initial concentration  $c_0$  and input parameters  $\mathbf{q}$ , produced concentration of species k at time t,  $c_{kt}$ . Nominating  $\mathbf{q}$  as independent variables and  $\mathbf{c}$  as dependent variables, ADIFOR then automatically produces a code that computes the desired sensitivity coefficients.

Carmichael et al. (1997) applied such a black-box approach to the STEM-II model using a quasisteady state approximation (QSSA) integrator. They showed that for a fixed-stepsize QSSA integrator this black-box approach delivered results that are comparable to those obtained by integrating the ODE's for  $S_{kl}$  that would be obtained by differentiation of Eq. (8). In general, though, the automatic differentiation of sophisticated numerical integrators requires some care. In particular, Eberhard and Bischof (1996) showed that for integrators with adaptive stepsize control (such as LSODE) some postprocessing may be required to recover the desired sensitivities. They observed that the *algorithmic* sensitivities (which are computed by AD) may reflect the impact of numerical artifacts such as stepsize control and showed the relation of these algorithmic sensitivities with the *analytic* sensitivities that are desired. In the case of LSODE, it turned out that the stepsize control did not affect the AD-computed derivatives as described in Eberhard and Bischof (1996). However, under cloudy conditions, we had to adjust the so-called "internal stepsize" parameter of LSODE to obtain the desired derivatives. We are continuing our studies on the interplay of AD and sophisticated numerical algorithms. So far initial work has been done on integrators (Carmichael et al., 1997; Eberhard and Bischof, 1996) and certain classes of nonlinear solvers (Griewank et al., 1993), but much remains to be done.

To compare sensitivities with dependent and independent variables of different orders of magnitude, the dimensionless sensitivities (i.e., normalized local sensitivities) is calculated:

$$\bar{S}_{kl} = \frac{q_l}{c_k} \frac{\partial c_k}{\partial q_l} = \frac{\partial \ln c_k}{\partial \ln q_l} \tag{10}$$

The sign of  $\bar{S}_{kl}$  gives the sense of response of  $c_k$  upon the relative variation of  $q_l$ . The positive values indicate that  $c_k$  increases with the increase in  $q_l$ , while the negative values mean a reversed change in  $c_k$  when increasing  $q_l$ . The magnitude of  $\bar{S}_{kl}$  gives the relative change in  $c_k$  due to one unit change (i.e., doubling its value) in  $q_l$ . For instance, a value of  $\bar{S}_{kl}$  =0.5 can be interpreted to mean that doubling  $q_l$  will lead to a 50% increase in concentration of species k.

Applying ADIFOR to MaTChM, we computed the sensitivities of 144 gas, aqueous and aerosol species concentrations with respect to 404 model parameters at every integration time step. The model parameters include 245 reaction rate constants; 134 mass transfer parameters; 21 species initial concentrations, and 4 physical parameters (temperature, R.H., cloud water content and droplet radius). While it is easy to compute derivatives with AD and recompute derivatives in response to changes in the code (we actually found a bug in the code of MaTChM through inspection of the ADIFOR-computed sensitivities), large-scale sensitivity analysis is still computationally expensive. For example, the undifferentiated model required 0.72MB of memory and 1.3 minutes on an IBM RS/6000 workstation with 128MB RAM running at 62.5 Mhz. On the same machine, the sensitivity analysis with respect to all 404 parameters required 197 MB of memory and roughly 10.6 hours CPU time. For every run, we computed 404 parameters x 144 concentrations x 60 time steps = 3,490,560 sensitivities. Using ADIFOR "out of the box," it was feasible to compute this large number of sensitivities with little human effort, and the time required to compute the derivatives mattered little in comparison to the effort required to interpret the 27.7 Mbytes of data produced by one run.

We note, however, that AD and the associativity of the chain rule allow for various ways of decreasing the computational cost, both with respect to memory and runtime. Derivatives can be computed in parallel (Bischof et al. 1994; Bischof and Wu 1997), cost can be reduced by techniques such as interface contraction (Hovland et al., 1997; Bischof and Haghighat, 1996), the number of derivatives can be reduced via "shaped" perturbations (Bischof et al., 1996c), or the understanding of the mathematical underpinnings of a particular algorithm may make it possible to avoid differentiation of parts of the code (Griewank et al., 1993; Carle and Fagan, 1996).

Given the sensitivities of individual species, the sensitivities of the ratio of the concentrations of the  $i^{th}$  species to the  $j^{th}$  species with respect to the arbitrary parameter  $q_l$  can be easily derived as the difference of the two individual sensitivities:

$$\bar{S}r_{ijl} = \frac{q_l}{(c_i/c_j)} \frac{\partial (c_i/c_j)}{\partial q_l} = \frac{q_l}{c_i} \frac{\partial c_i}{\partial q_l} - \frac{q_l}{c_j} \frac{\partial c_j}{\partial q_l} = \bar{S}_{il} - \bar{S}_{jl}$$
(11)

The sensitivity of lumped concentrations of two (or more) species can also be calculated as the sum of products of the concentration fraction and the sensitivity coefficient of each species:

$$\overline{S}_{Sijl} = \frac{q_l}{(c_i + c_j)} \frac{\partial (c_i + c_j)}{\partial q_l} = \frac{c_i}{(c_i + c_j)} \overline{S}_{il} + \frac{c_j}{(c_i + c_j)} \overline{S}_{jl}$$
(12)

Using Eq. (11) and (12), we calculated the sensitivities of several lumped species (e.g.,  $NO_x$ ,  $NO_z$ ,  $R_xO_y$ , defined in Table 1) and photochemical indicators (e.g.,  $NO_y$ ,  $O_3/NO_z$ , and  $H_2O_2/HNO_3$ ) with respect to all model parameters under all conditions. Instantaneous sensitivities at selected time steps and the sensitivities averaged during the last hour simulation along with base simulation results are analyzed and discussed below.

### 4. Results and Discussions

#### 4.1 The Base Simulations

## 4.1.1 Heterogeneous Effects on Gas-Phase Species

Table 5 shows the average concentrations and percent changes in concentrations of 12 major gasphase species and total sulfate and nitrate formed in all phases during the last hour simulation under various conditions. The heterogeneous processes associated with clouds and aerosols can affect these species in a variety of ways, either increasing or reducing their concentrations. The changes in species concentrations range from a few percent to several times, depending on properties of species (e.g., gas-phase diffusivity, interfacial mass accommodation coefficient or uptake coefficient, effective solubility, diffusivity, and

reactivity in the aqueous-phase) and properties of pre-existing particles (e.g., liquid water content and size distribution).

Under cloudy conditions, HNO<sub>3</sub>, HCHO, HO<sub>2</sub>, and H<sub>2</sub>O<sub>2</sub> experience significant reduction (10.4-100%) in their gas-phase concentrations due to their high solubilities (with Henry's law constants greater than 100 M atm<sup>-1</sup>). Although OH, SO<sub>2</sub>, and H<sub>2</sub>SO<sub>4</sub> have relatively small Henry's law constants, these species can undergo rapid aqueous-phase reactions and/or dissociations, which greatly enhances their effective solubility and results in large concentrations decreases (6.8-100%). Under aerosol conditions, various species undergo irreversible reactions on aerosols with a pseudo first-order rate ranging from 1 x 10<sup>-10</sup> to 1 x10<sup>-2</sup> s<sup>-1</sup>, which depends on gas-phase diffusivity, pre-existing aerosol surface and individual species uptake coefficient. When the surface areas are greater than 1 x10<sup>2</sup>  $\mu$ m<sup>2</sup> cm<sup>-3</sup> (i.e., Rural), aerosols are found to have appreciable effects (decrease by 6.6- 43.9%) on many species including HNO<sub>3</sub>, PAN, H<sub>2</sub>SO<sub>4</sub>, HCHO, OH, HO<sub>2</sub>, and H<sub>2</sub>O<sub>2</sub>. They become very effective scavengers when the available surface areas are greater than 1 x10<sup>3</sup>  $\mu$ m<sup>2</sup> cm<sup>-3</sup> (i.e., Urban and Heavily-polluted), with decreases in many species concentrations comparable or even larger than those under cloudy conditions.

In the presence of both clouds and aerosols, heterogeneous uptake and chemistry on both surfaces occur, competing for the available condensing gases. The combined heterogeneous effects are a nonlinear function of their individual effects. The magnitude of these effects is determined by the scavenging efficiencies of clouds and aerosols, which vary with the initial gas-phase concentrations, the assumed cloud parameters and aerosol size distributions. In the Marine and Remote atmospheres, cloud processes control losses of almost all species, because the total surface area of cloud droplets is 1.2 x 10<sup>5</sup> μm<sup>2</sup> cm<sup>-3</sup>, which is at least four orders of magnitude higher than the total aerosol surface areas. Aerosol surface uptake becomes increasingly important with increasing surface areas. Under the Heavily-polluted and the Urban conditions, the large aerosol surface areas result in rapid heterogeneous loss rates for many condensing species. Consequently, aerosols serve as a very effective sink for many species including OH, HO<sub>2</sub>, HCHO, PAN, and O<sub>3</sub>, while both clouds and aerosols contribute significantly to the loss of HNO<sub>3</sub>, SO<sub>2</sub>, H<sub>2</sub>SO<sub>4</sub>, and H<sub>2</sub>O<sub>2</sub>. The effect of aerosols can be comparable or even overwhelming to that of clouds. For instance, under the Heavily-polluted conditions, more than 80% of decrease in O<sub>3</sub> and PAN concentrations is due to aerosols surface uptake processes.

As a consequence of the decreases in concentrations of many gaseous precursors and reactive radicals, the predicted concentrations of  $O_3$  and PAN are consistently reduced in the presence of clouds and/or aerosols. Figure 1 shows concentrations of  $O_3$  and PAN as a function of time under various polluted conditions. The decreases in  $O_3$  concentrations at the end of the 2-hour simulation range from 0.6-5.1 ppb (~3-9%), 0.2-24.2 ppb (~0.3-27%) and 0.6-24.5 ppb (~3-26.8%) due to cloud chemistry, aerosol surface uptake, and the combined cloud and aerosol chemistry, respectively. While the magnitude of cloud effect on  $O_3$  is similar under various conditions due to constant cloud water content and droplet size, the aerosol surface area and  $O_3$  formation are anti-correlated, i.e., the magnitude of  $O_3$  reduction appreciably increases as the air becomes increasingly polluted (i.e., the surface areas become larger). Under moderately-polluted and relatively clean conditions, the decreases in  $O_3$  due to aerosol surface reactions are negligible (< 2%). The aerosol surface uptake processes become important and dominate the decreases in  $O_3$  concentrations under

the Urban and the Heavily-polluted conditions. The relative changes in PAN concentrations under various conditions are even larger, with decreases of 0.1-0.7 ppb (~8-69.2%), 0.003-1.7 ppb (~2-38%) and 0.15-2.1 ppb (33-69.2%) at the end of simulation due to cloud chemistry, aerosol surface uptake and their combination, respectively.

Clouds and aerosols are not always a sink for gas-phase species. Their presences actually help increase gas-phase budgets of some insoluble or less soluble reactive nitrogen and organic species including NO, NO<sub>2</sub>, PAR, OLE, ETH, TOL, XYL and ISOP. Since the reactions of these species with many oxidants and radicals are usually the major sinks, the decrease in the concentrations of oxidants and radicals reduced these sinks. For example, lower OH and HO<sub>2</sub> can lower the depletion rates of NO, PAR, OLE, ETH, TOL, XYL, and ISOP via reactions G28, G52, G57, G61, G63, G69 and G76, respectively. As a result, these species remain in the gas-phase with higher concentrations when there are clouds and/or aerosols. The effects of clouds and aerosols are complicated for NO<sub>2</sub>, with a decreased concentration under the Remote and Heavily-polluted conditions and an increased one under other conditions, reflecting a competition between its sink and source reactions. In addition, the heterogeneous oxidation of SO<sub>2</sub> and NO<sub>x</sub> associated with clouds and aerosols is shown to be a dominant pathway, resulting a higher total sulfate formation under all cloud and/or aerosol conditions and a higher nitrate formation under the Heavily-polluted conditions as compared to those in clear air.

Dentener and Crutzen (1993) reported a net decrease in yearly-averaged concentrations of  $NO_x$  (by 50%),  $O_3$  (by 25%) and OH (by 30%) due to heterogeneous reactions of  $NO_3$  and  $N_2O_5$  on aerosols using a 3-D global model. While consistent results are found for  $O_3$  and OH, our results also show the concentrations of  $NO_x$  could be higher (rather than lower) when aerosol chemistry is included. The different aerosol effects on  $NO_x$  can be attributed to two reasons. First, the uptake coefficients that we use for  $NO_3$  and  $N_2O_5$  are 1 x  $10^{-3}$  and 5 x  $10^{-3}$ , respectively (Thomas et al., 1989; Kirchner et al., 1990), which are lower limits of available measurements. Dentener and Crutzen adopted the value of 0.1 for both species, which is the upper limit of measurements. While these two studies provide the variation range of the observation, the rate of denitrification due to  $NO_3$  and  $N_2O_5$  reactions on aerosols is several orders of magnitude faster in their work. Secondly, Dentener and Crutzen neglected other possible aerosol surface reactions such as uptake of OH,  $HO_2$  and  $H_2O_2$  in their calculations. OH levels are likely overpredicted when these processes are ignored. The lowered OH levels predicted in this work can further reduce the rate of  $NO_x$  sinks (e.g, G26 and G28), resulting in higher gaseous  $NO_x$ .

## 4.1.2 Heterogeneous Effects on Photochemical Indicators

Scientific understanding of the relationships between  $O_3$  and its precursors  $NO_x$  and ROG is essential in the development of more effective  $O_3$  abatement strategies. A fundamental factor that leads to difficulties and ineffectiveness in attainment of  $O_3$  is that  $O_3$  is not directly emitted but is formed by a complex set of photochemical reactions involving nonlinear interactions among  $NO_x$ , ROG, and CO. In the past, evaluations of  $O_3$ - $NO_x$ -ROG sensitivity have largely relied on 3-D photochemical model simulations, which are highly dependent upon model assumptions and thus possibly lead to inaccurate predicted sensitivity.

Recent studies have established the use of measurable photochemical indicators in determining the  $O_3$ -NO<sub>x</sub>-ROG sensitivity in order to effectively control  $O_3$  at non-attainment regions. This approach is based on direct measurements of individual species or species ratios that consistently assume different values under  $NO_x$ - and ROG-sensitive conditions. Milford et al. (1994) have found that  $O_3$  is ROG-sensitive in regions where afternoon  $NO_y$  concentrations are higher than a threshold value ranging from 10-25 ppb, whereas it becomes  $NO_x$ -sensitive with  $NO_y$  concentrations lower than the threshold. Sillman (1995) has linked ROG-sensitivity chemistry to afternoon conditions with  $NO_y$ >20 ppb,  $O_3/NO_z$  < 7, HCHO/ $NO_y$  < 0.28, and  $H_2O_2/HNO_3$  < 0.4, lower  $NO_y$  and higher ratios correspond to  $NO_x$ -sensitive  $O_3$ . Other indicators include  $O_3/NO_y$ ,  $O_3/HNO_3$ ,  $H_2O_2/NO_y$  (Trainer et al., 1993; Sillman, 1995) and  $(HNO_3/H_2O_2)$  /  $(NO_x/ROG)$  (Stockwell, 1986). All these indicators and corresponding threshold values are established under the clear air conditions. The theoretical rational for use of these species and ratios as indicators for  $O_3$ -NO<sub>x</sub>-ROG sensitivity is described in several literatures (e.g., Milford et al., 1994; Sillman, 1995).

The calculated indicators under various polluted conditions are shown in Table 6. The relative changes represent the perturbation in the indicators due to clouds and/or aerosols relative to the values under the clear air conditions. According to the aforementioned threshold values of indicators, O<sub>3</sub> is ROGsensitive and NO<sub>x</sub>-sensitive under the Heavily-polluted and the Rural clear air conditions, respectively. O<sub>3</sub> chemistry exhibits a transition between the two regimes under the Urban conditions, with  $10 < NO_y < 20~\text{ppb}$ but values of O<sub>3</sub>/NO<sub>z</sub>, HCHO/NO<sub>v</sub> and H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub> higher than the thresholds. While cloud and aerosol chemistry can always increase O<sub>3</sub>/NO<sub>z</sub> and O<sub>3</sub>/HNO<sub>3</sub>, decrease NO<sub>v</sub>, H<sub>2</sub>O<sub>2</sub>/NO<sub>v</sub>, HCHO/NO<sub>v</sub>, and (HNO<sub>3</sub>/H<sub>2</sub>O<sub>2</sub>)/(NO<sub>x</sub>/ROG) under all modeled conditions, it can cause either an increase or a decrease in O<sub>3</sub>/NO<sub>v</sub> and H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub>. Some changes are so substantial that they can cause a controversy or even failure of the above criteria developed for determination of O<sub>3</sub>-NO<sub>x</sub>-ROG sensitivity. For example, the value of O<sub>3</sub>/NO<sub>2</sub> under the Heavily-polluted conditions changes from 6.48 to 14.1 and 13.2 in the presence of clouds and aerosols, respectively, which exceeds the range of ROG-sensitive regime. Other indicators NO<sub>v</sub>, HCHO/NO<sub>v</sub> and H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub> still fall in the ROG-sensitivie regime. Exclusion of heterogeneous chemistry could cause the incorrect model predictions of indicators and the inaccurate determination of threshold values for different chemical regimes. Therefore, the established criterias of indicators under clear air conditions are not applicable to regions where the large cloud coverage or high aerosol loading are frequently observed.

## 4.2 Sensitivity Analysis

The base simulation results show that heterogeneous chemistry can significantly change concentrations of many important gas-phase species and reduce the total tropospheric oxidizing capability (TOC) (i.e., the total burden of principal oxidants such as O<sub>3</sub>, H<sub>2</sub>O<sub>2</sub>, and OH radicals) under typical atmospheric conditions. To further study the correlation of O<sub>3</sub> and its precursors and to evaluate the relative importance of the homogeneous and heterogeneous pathways, we conduct several sets of sensitivity analysis for the mixed-phase chemical mechanism by excluding or including heterogeneous chemistry under the five

conditions using automatic differentiation ADIFOR tool. The results are presented and discussed in the following sections.

## 4.2.1 Gas-Phase Chemistry

#### 4.2.1.1 Sensitivity to Gas-Phase Reaction Rate Constants

The 30 most influential reactions for O<sub>3</sub> concentrations under the five conditions are ranked along with the 1-hr average sensitivities of O<sub>3</sub> to these reaction rate constants in Table 7. These reactions could have either positive or negative influences depending on the signs of their sensitivities. O<sub>3</sub> predictions are more sensitive (by a factor of a few to several hundreds) to reaction rate constants under various polluted conditions (i.e., Rural, Urban and Heavily-polluted) than relatively clean conditions (i.e., Marine and Remote) for one unit change in the rate constants. For instance, when the rate constant of R1 doubles, the predicted O<sub>3</sub> concentration will increase by 1.86%, 4.13% and 27.7 % under the Remote, Rural and Heavily-polluted conditions, respectively. This is because O<sub>3</sub> formation rate in the lower atmosphere is proportional to the TOC. The TOC is much higher under polluted conditions, resulting in higher reaction rates of most species. Higher sensitivities reflect higher influence of these reactions and larger model responses to uncertainties in the rate constants.

Under the Remote and Marine conditions, the most important reactions contributing to O<sub>3</sub> production are NO+HO<sub>2</sub> (G28), NO<sub>2</sub>+hv (G1), CH<sub>4</sub>+OH (G116), NO+CH<sub>3</sub>O<sub>2</sub> (G117), H<sub>2</sub>O<sub>2</sub>+hv (G34), CO+OH (G36), HCHO+hv (G38), C<sub>2</sub>O<sub>3</sub> + NO (G46), the photolysis of O<sub>3</sub> (G9) and the subsequent reaction of O(<sup>1</sup>D)+H<sub>2</sub>O (G11). G28 and G117 are major pathways for conversion of NO to NO<sub>2</sub>. Followed by G1, they generate O(3p), which directly produces O<sub>3</sub>. The photolysis of H<sub>2</sub>O<sub>2</sub> (G34), HCHO (G38) and O<sub>3</sub> (G9) are the major sources of OH and HO<sub>2</sub>. A higher rate constant of G9 decreases O<sub>3</sub> but increases O(<sup>1</sup>D), which contributes to O<sub>3</sub> production via generation of OH (G11). The increase in O<sub>3</sub> via G11 compensates the decrease via G9. Thus, G9 has a net positive impact on O<sub>3</sub>. OH can be converted to HO<sub>2</sub> and CH<sub>3</sub>O<sub>2</sub> by reacting with CO (G36) and CH<sub>4</sub> (G116), respectively. These radicals then convert NO to NO<sub>2</sub> via G28, G117 and G46. The generation and inter-conversion of these radicals thus increase O<sub>3</sub> by increasing the TOC. Compared to the Remote conditions, O<sub>3</sub> formation under the Marine conditions is more affected by the photolysis of HCHO (G38) by one order of magnitude, due to a much higher initial HCHO. The reactions contributing to O<sub>3</sub> destruction include NO+O<sub>3</sub> (G3), HO<sub>2</sub>+CH<sub>3</sub>O<sub>2</sub> (G118), NO<sub>2</sub>+OH (G26), 2HO<sub>2</sub>+H<sub>2</sub>O (G33), HO<sub>2</sub>+HO<sub>2</sub> (G32),  $O(^{1}D) \rightarrow O(^{3}p)$  (G10),  $O_{3}+HO_{2}$  (G13),  $O_{3}+OH$  (G12),  $ALD_{2}+OH$  (G43), and  $C_{2}O_{3}+NO_{2}$  (G47).  $O_3+NO$  (G3) is the predominant destruction pathway for  $O_3$ . G10 reduces  $O_3$  via reducing  $O(^1D)$ , a major source of OH. Representing the major sink for OH, HO<sub>2</sub>, and CH<sub>3</sub>O<sub>2</sub>, the formation of peroxides (G118, G33 and G32), HNO<sub>3</sub> (G26) and PAN (G47 following G43) reduces the TOC thus decreases O<sub>3</sub>. O<sub>3</sub> can also be directly destroyed by OH (G12) and HO<sub>2</sub> (G13).

Under various polluted conditions, G28, G32, G33, G116-118 become less important or unimportant because of the relatively lower HO<sub>2</sub> and CH<sub>3</sub>O<sub>2</sub>. O<sub>3</sub> predictions are highly sensitive to ROG chemistry. This is consistent with the results from photochemical indicator calculations in section 4.1.2. HCHO+hv

(G38), ALD<sub>2</sub>+OH (G43), OLE+OH (G57) become the most important positively-influential reactions under the Urban and the Heavily-polluted conditions, with their effects comparable or even higher than other important inorganic reactions such as G1 and G3. Other important positively-influential ROG reactions include reactions of C<sub>2</sub>O<sub>3</sub> with NO and NO<sub>2</sub> (G46 and G47), photolysis of ALD<sub>2</sub> and MGLY (G45 and G74), OLE+O<sub>3</sub> (G58), NO+TO<sub>2</sub> (G64) and reactions of HCHO, PAR, ETH, TOL, XYL and ISOP with OH (G37, G52, G61, G63, G69, and G76). A higher rate constant of G58 directly decreases O<sub>3</sub> but simultaneously increases odd hydrogen R<sub>x</sub>O<sub>y</sub> and O<sub>3</sub> precursors HCHO and ALD<sub>2</sub>, which eventually give rise to O<sub>3</sub> via G1 and G2. The increase in O<sub>3</sub> due to these pathways suppresses the direct decrease, thus G58 shows a net positive impact. Other ROG reactions (i.e., G37, G45, G46, G52, G57, G61, G63, G64, G69, and G74) contribute to O<sub>3</sub> formation by producing HCHO, ALD<sub>2</sub>, HO<sub>2</sub> and RO<sub>2</sub>. G43 and G47 are dominant negatively-influential ROG reactions. G39 also contributes to O<sub>3</sub> destruction by competing with the positively-influential reaction G38 and consuming HCHO. These results are consistent with the sensitivity analyses of other gas-phase mechanisms such as RADM2, CBM-IV, LLM and STEM-II under smog chamber and urban plume conditions by Milford et al. (1992), Gao et al. (1995) and Carmichael et al. (1997).

Photochemical indicators are mainly discovered based on theoretical rational and case studies using 3-D models, therefore their applications involve large uncertainties arising from model assumptions or omissions. While being verified by growing observational dataset, the sensitivity of these model-based indicators to model parameters have not been fully explored. Our sensitivity analysis shows that  $NO_y$  and  $O_3/NO_y$  are relatively insensitive ( $\overline{S}_{Rl} < 0.01$ ) to most model parameters (e.g., reaction rate constants), whereas other indicators involving  $H_2O_2$ ,  $HNO_3$ , HCHO and  $NO_z$  are highly sensitive to changes in many model parameters. Figure 2 shows the 1-hr average sensitivities of  $O_3/NO_z$  and  $H_2O_2/HNO_3$  with respect to gas-phase reaction rate constants under various polluted conditions. Both  $O_3/NO_z$  and  $H_2O_2/HNO_3$  are much more sensitive to these rate constants than  $O_3$  itself, with their sensitivities as large as 0.6.

While the most influential reactions for these indicators are almost the same as those for O<sub>3</sub>, their relative impacts are different both in signs and amplitudes. For example, the photolysis of HCHO (G38) has a positive impact on O<sub>3</sub>, but a negative impact on O<sub>3</sub>/NO<sub>z</sub> and H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub>. While a higher rate constant of G38 increases O<sub>3</sub> and H<sub>2</sub>O<sub>2</sub> formation via generation of HO<sub>2</sub>, it also increases the denominator NO<sub>z</sub> and HNO<sub>3</sub>. The net effect is the decrease in both O<sub>3</sub>/NO<sub>z</sub> and H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub>. Such a decrease can be up to 10-55% when doubling the reaction rate constant of G38. Similar to O<sub>3</sub>, the indicators generally exhibit higher sensitivities on basic O<sub>3</sub>-NO<sub>x</sub>-CO reactions (G9-G11, G26, G28, G29, G32-34, and G36) under the Rural conditions, and higher (by a factor of 2-30) sensitivities on various ROG reactions (G38, G39, G43, G45-G47, G52, G57, G58, G61, G69, and G74) under the Urban and the Heavily-polluted conditions. This coincides with the nature of the indicators as good markers for the O<sub>3</sub>-NO<sub>x</sub>-ROG sensitivity. For instance, the sensitivities of O<sub>3</sub>/NO<sub>z</sub> and O<sub>3</sub> to G38, G45, and G58 under the Heavily-polluted conditions are higher than those under the Rural conditions by a factor of 4.0, 16, and 29, and a factor of 18, 68, and 168, respectively, indicating that ROG chemistry is dominant under the Heavily-polluted conditions. Lower

sensitivities to ROG reactions and higher sensitivities to  $O_3$ -NO<sub>x</sub>-CO reactions under the Rural conditions indicate a NO<sub>x</sub>-dominant chemistry.

## 4.2.1.2 Sensitivity to Species Initial Concentrations

The O<sub>3</sub> predictions are also sensitive to species initial concentrations. The initial O<sub>3</sub> always has the predominant positive impact on the predicted O<sub>3</sub>. Doubling initial O<sub>3</sub> will result in an increase of 73-100% in predicted O<sub>3</sub> under various conditions. Figure 3(a) shows the sensitivity of O<sub>3</sub> with respect to initial species concentrations under the five conditions at the end of the 2-hr simulation. The O<sub>3</sub>-precursor relations are different at various conditions. Under relatively-clean conditions, O<sub>3</sub> production increases with increasing NO<sub>x</sub> but is insensitive to ROG (except for HCHO and ALD<sub>2</sub> under the Marine conditions). At various polluted conditions, O<sub>3</sub> formation increases rapidly with increasing ROG (except for ALD<sub>2</sub>) and decreases with increasing NO<sub>x</sub>. The sensitivities of O<sub>3</sub> to positively-influential ROG species increase as the air becomes increasingly polluted. Other O<sub>3</sub> precursors such as H<sub>2</sub>O<sub>2</sub>, CO, and CH<sub>4</sub> have relatively large impact on O<sub>3</sub> production under the Remote, Marine and Rural conditions. Being a major source of OH, H<sub>2</sub>O<sub>2</sub> helps produce O<sub>3</sub> under these conditions. CO and CH<sub>4</sub> always tend to increase O<sub>3</sub> via G36 and G116 followed by G28, G1 and G2.

Since O<sub>3</sub> production rate is proportional to reactions of ROG with OH radicals under most atmospheric conditions (Sillman, 1995), the different O<sub>3</sub>-precursor relations under various conditions can be interpreted relative importance of reactions of odd by hydrogen,  $R_xO_v$ (=OH+HO<sub>2</sub>+CH<sub>3</sub>O<sub>2</sub>+ETHP+C<sub>2</sub>O<sub>3</sub>+ROR+TO<sub>2</sub>+CRO, see Table 1), as shown in Table 8. Under the Remote and Marine conditions, formation of peroxides via HO<sub>2</sub>+HO<sub>2</sub> (G32), 2HO<sub>2</sub>+H<sub>2</sub>O (G33) and HO<sub>2</sub>+CH<sub>3</sub>O<sub>2</sub> (G118) are the major sinks for  $R_xO_y$ . The fate of OH is mainly governed by the inter-conversion of OH, HO<sub>2</sub>, and CH<sub>3</sub>O<sub>2</sub> via NO+HO<sub>2</sub> (G28), CH<sub>4</sub>+OH (G117), CO+OH (G36), and HCHO+OH (G37). OH decreases with a higher HCHO (due to G37) under both conditions and other ROG species including ALD<sub>2</sub>, PAR, OLE, C<sub>2</sub>H<sub>6</sub>, ETH and TOL (due to G37, G43, G47, G51, G52, G57 and G61) under the Marine Thus, O<sub>3</sub> production is insensitive to most ROG (only HCHO exists under the Remote conditions), because increases in ROG coincide with decreases in OH. As a predominant pathway for O<sub>3</sub> production, the conversion of NO by HO<sub>2</sub> (G28) results in a higher O<sub>3</sub> followed by G1 and G2 (see Table 7). NO+HO<sub>2</sub> (G28) and NO<sub>2</sub>+hv (G1) compete with NO+O<sub>3</sub> (G3) and NO<sub>2</sub>+OH (G26), with less NO<sub>x</sub> available for the latter reactions. In addition, OH increases with increasing NO<sub>x</sub> (due to G28). Thus, a higher NO<sub>x</sub> always causes a higher O<sub>3</sub> under both conditions.

Under various polluted conditions, the reactions of  $NO_2$ +hv (G1),  $NO_2$ +O<sub>3</sub> (G3),  $NO_2$ +OH (G26), HCHO+hv (G38), ALD<sub>2</sub>+OH (G43), and  $NO_2$ +C<sub>2</sub>O<sub>3</sub> (G47) are the overwhelming pathways affecting O<sub>3</sub> production. Both NO+HO<sub>2</sub> (G28) and NO<sub>2</sub>+OH (G26) are the predominant pathways for  $R_xO_y$ . While an increase in NO consumes more O<sub>3</sub> and HO<sub>2</sub> via G3 and G28, a higher NO<sub>2</sub> results in a lower OH (due to G26). Therefore, an increase in  $NO_x$  leads to a lower O<sub>3</sub>. Increases in individual ROG species either increase or decrease OH, but the net overall effect is a slight increase in OH, reflecting that ROG reactions

could be a source of  $R_xO_y$ . Thus,  $O_3$  increases with increases in most ROG species. When presence,  $ALD_2$  always has a negative influence on  $O_3$  (up to 11% decrease) because it acts as a sink for  $NO_x$  via G43 and G47. G43 generates  $C_2O_3$ , which either reduce  $NO_2$  through G47 or convert NO to  $NO_2$  via G46. Being a sink for  $NO_2$ , G47 is relatively more important to the source G46. Correspondingly,  $O_3$  decreases with higher rate constants of G43 and G47 but increases with a higher rate constant of G46 (see Table 7), causing the anti-correlation between  $O_3$  and the initial  $ALD_2$ . The negative impact of  $ALD_2$  on  $O_3$  formation has also been reported by Carmichael et al. (1997) under similar conditions, by Carter and Atkinson (1989) under low  $NO_x$  conditions and by Milford et al. (1992) under the smog chamber conditions with higher ROG to  $NO_x$  ratios.

Figure 3 (b) through (d) show the 1-hr average sensitivities of NO<sub>v</sub>, O<sub>3</sub>/NO<sub>z</sub> and H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub> to species initial concentrations under various polluted conditions. These indicators are affected in a variety of ways. NO<sub>v</sub> is predominantly correlated to initial NO<sub>x</sub> and almost insensitive to initial concentrations of other species. O<sub>3</sub>/NO<sub>z</sub> is anti-correlated to initial H<sub>2</sub>O<sub>2</sub>, NO, HCHO, ALD<sub>2</sub>, OLE, TOL, XYL and ISOP, and positively-correlated with initial O<sub>3</sub>, PAR, CO and CH<sub>4</sub>. The impacts of NO<sub>2</sub> and ETH are complicated and can be either way. For H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub>, the positively-influential species include H<sub>2</sub>O<sub>2</sub>, ALD<sub>2</sub>, PAR, CO and CH<sub>4</sub>, and the negatively-influential species are O<sub>3</sub>, NO<sub>x</sub>, HCHO and XYL. The impacts of OLE, ETH and TOL can be either negative or positive. These relations are much more complicated relative to O<sub>3</sub>precursor relations. The interpretation requires an understanding of the chemical behaviors of the individual indicator species. Although an increase in various ROG species (other than ALD<sub>2</sub>) can always increase O<sub>3</sub> production, only a few of them (e.g., PAR) can increase O<sub>3</sub>/NO<sub>z</sub>. Most ROG species negatively affect the indicator because they simultaneously increase the denominator NO<sub>z</sub> to the extent of exceeding the increase in O<sub>3</sub>. Whether the species have positive or negative impacts depends on the relative importance of their reactions and the signs of the predominant reactions. For instance, G1, G26 and G28 are important reactions for NO<sub>2</sub>. Under the Rural and Urban conditions G26 and G28 are the predominant reactions and both have a Under the Heavily-polluted conditions, G1 is predominant and has a positive sign. Correspondingly, O<sub>3</sub>/NO<sub>2</sub> and NO<sub>2</sub> are positively-correlated under the Heavily-polluted conditions but anticorrelated under the Rural and Urban conditions.

## 4.2.1.3 Sensitivity to Temperature and Relative Humidity

Most species are very sensitive to changes in temperature and relative humidity (R.H.) under all modeled conditions, as shown in Table 9. Under all conditions, a higher temperature always results in higher HNO<sub>3</sub>,  $H_2SO_4$ , OH,  $HO_2$ , PAN and  $O_3$  due to an increase in their formation rates and lower  $SO_2$  due to an increase in its destruction rate. A higher temperature will decrease  $H_2O_2$  and increase  $O_3$ ,  $NO_2$  and  $HNO_3$ , with the net effects of increasing  $NO_y$  and decreasing the other two indicators.  $O_3$  (under relatively-clean conditions),  $SO_2$  (under all conditions), and  $NO_y$  (under polluted conditions) are less sensitive to changes in temperature, with their sensitivities at least one order of magnitude lower than those of other species and indicators. For example, an increase of 2.8 K in temperature will increase  $O_3$  by 1.26% but increase  $H_2SO_4$ 

by 16.7% and decrease  $O_3/NO_z$  by 12.8% under the Rural conditions. While  $SO_2$ ,  $O_3$  and  $NO_y$  are also less sensitive to R.H., other species and indicators exhibit much higher sensitivities to R.H., with values greater than 1 x  $10^{-1}$ . This is because the fates of these species and indicators are governed by OH, which is mainly produced through reaction of  $O(^1D)+H_2O$  (G11) and highly depends on the ambient concentration of water vapor.

## 4.2.2 Aqueous-Phase Chemistry

## 4.2.2.1 Sensitivity to Aqueous-Phase Kinetic Reactions

The 25 most influential aqueous-phase reactions on gas-phase O<sub>3</sub> concentrations under various cloudy conditions are shown in Table 10. Various aqueous-phase reactions play an important role under relatively clean (i.e., Remote and Marine) and moderately-polluted (i.e., Rural) conditions, with their sensitivities comparable to other dominant gas-phase reactions. The sensitivities of O<sub>3</sub> formation to aqueous-phase reaction rate constants are at least one order of magnitude smaller than other dominant gas-phase reactions under the Urban and Heavily-polluted conditions. Most aqueous-phase reactions have negative effects on gas-phase O<sub>3</sub> formation throughout the simulation time because these reactions accelerate the rates of scavenging of various gas-phase species including many O<sub>3</sub>-precursors. As a result, the formation rate of O<sub>3</sub> drops from 0.3 to 0.0 ppb hr<sup>-1</sup>, 2.25 to 0.6 ppb hr<sup>-1</sup>, 2.2 to 0.75 ppb hr<sup>-1</sup>, 9.07 to 5.75 ppb hr<sup>-1</sup>, and 5.88 to 3.05 ppb hr<sup>-1</sup> under cloudy Remote, Marine, Rural, Urban and Heavily-polluted conditions, respectively, as compared to the corresponding clear air conditions.

The major aqueous-phase reactions include HO<sub>2</sub> + O<sub>2</sub>- (L7), O<sub>3</sub> + O<sub>2</sub>- L(13), SO<sub>3</sub><sup>2</sup>-+O<sub>3</sub> (L74), CH<sub>2</sub>(OH)<sub>2</sub>+OH (L50), HSO<sub>3</sub>-+H<sub>2</sub>O<sub>2</sub> (L75), HO<sub>2</sub>+HO<sub>2</sub> (L6), and S(IV)+HO<sub>2</sub> (L105) under all conditions; H<sub>2</sub>O<sub>2</sub>+O<sub>3</sub> (L16) under relatively clean conditions; and SO<sub>5</sub>-+O<sub>2</sub>- (L82), 2NO<sub>2</sub>+HSO<sub>3</sub>- (L109), and HCHO+ SO<sub>3</sub><sup>2</sup>- (L112) under polluted conditions. Reactions L6, L7, L13, L16, L74, and L112 act as aqueous-phase sinks for HO<sub>2</sub>, H<sub>2</sub>O<sub>2</sub>, O<sub>3</sub> and HCHO, thus decrease O<sub>3</sub> formation. Other reactions such as L50, L75, L82, L105, L109 can affect O<sub>3</sub> in a variety of ways. For example, L75 has a negative impact under polluted conditions but positive under relatively clean conditions. L50 shows an opposite impact. Two of these reactions

and L13 
$$O_3 + O_2^- \xrightarrow{H2O} O_1 + O_2 + O_3 + O_2^-$$
 OH +  $2O_2 + O_3 + O_3 + O_2^- \longrightarrow O_3 + O_3$ 

are the predominant aqueous-phase reactions under all conditions except the Heavily-polluted conditions. Followed the dissolution equilibrium of  $HO_2$  and  $O_3$  between gas and aqueous-phases, the self-reaction of dissolved  $HO_2$  (L7) destroys gas and aqueous-phase  $HO_2$  and L13 reduces not only gas and aqueous-phase  $HO_3$ , but more importantly gas and aqueous-phase  $HO_3$ . Being the dominant aqueous-phase sinks for  $HO_2$  and  $HO_3$ , impact of L7 and L13 is comparable to or even larger than their major gas-phase sink reactions such

as  $O_3+NO$  (G3),  $NO+HO_2$  (G28) and  $NO_2+OH$  (G26). These two reactions significantly reduce the gasphase concentration of  $HO_2$  and  $O_3$ , thus exhibit a negative impact on  $O_3$  formation. These results are in good agreement with the work of Walcek et al. (1997), in which they found a 30-90% reduction in local  $O_3$ formation rates due predominantly to the two aqueous-phase reactions L7 and L13 when excluding other trace metal reactions with dissolved  $HO_2$  under cloudy conditions.

Two aqueous-phase reactions are predominant under the Heavily-polluted conditions:

L109 
$$2NO_2+HSO_3^- \xrightarrow{H^2O} SO_4^{2-} + 3H^+ + 2NO_2^-$$
  
L75  $H_2O_2+HSO_3^- \xrightarrow{H^+} S(VI) + H_2O$ 

Although the rate of L109 is just 0.02%  $NO_2$ /hour, it results in a lifetime of  $NO_2$  in the aqueous-phase in the order of 1 s due to the low solubility of  $NO_2$ , thus  $NO_2$  plays an important role in the aqueous-phase chemistry when its concentrations are high. This reaction has a positive impact on  $O_3$  formation by consuming  $HSO_3$ - thus lowering the rates of other negatively-influential S(IV) conversion reactions such as  $H_2O_2+HSO_3$ - (L75),  $S(IV)+HO_2$  (L105),  $HCHO+SO_3$ <sup>2</sup>- (L112) and  $O_3+SO_3$ <sup>2</sup>- (L74). The secondary important reactions under polluted conditions are:

These two reactions reduce HO<sub>2</sub> and HCHO, thus exhibit negative impact on O<sub>3</sub> formation.

Pandis and Seinfeld (1989) identified aqueous-phase reactions important to major aqueous-phase species through a comprehensive sensitivity analysis using the direct decoupled method under the condition representative of a daytime cloudy environment in the northeastern United States. However, no gas-phase reactions were included in their simulations. Neglecting gas-phase reactions could lead to significant difference in chemical behaviors of species, especially when rates of gas-phase reactions are comparable or even higher than corresponding aqueous-phase reactions. Although a strict comparison between our results and their work is impossible because of differences in initial conditions, some reaction rate constants, and mass transfer treatment, Table 11 summarizes the 1-hr average sensitivities of aqueous-phase S(IV) and S(VI) under the five modeled conditions for a synoptic comparison.

As mentioned in Section 4.2.1.2, the formation of  $H_2O_2$  via G32 and G33 in the gas-phase is the major sink for odd hydrogen  $R_xO_y$  under the Remote and Marine conditions. Gaseous  $H_2O_2$  can be rapidly scavenged into cloud droplets due to its high solubility, leading to high aqueous-phase  $H_2O_2$  concentrations. With continuous sources (due to G32 and G33) and slow gas-phase sinks (due to G34 and G35), the high aqueous-phase  $H_2O_2$  can be maintained throughout the simulation time. The reaction of  $S(IV)+H_2O_2$  (L75) is thus the dominant pathway for oxidation of S(IV) and formation of S(VI) in the aqueous-phase and has a negative effect on aqueous S(IV) and a positive effect on S(VI) throughout the simulation. Other important pathways for conversion of S(IV) to S(VI) include oxidation of S(IV) by S(IV) include oxidation of S(IV) by S(IV) include oxidation of S(IV) include oxidation of S(IV) include oxidation of S(IV)

and  $Mn^{2+}$ ) (L77) and  $O_2^-$  (L106); oxidation of  $HSO_3^-$  by OH (L79) and  $CH_3OOH$  (L103); and oxidation of  $SO_3^{2-}$  by HCHO (L112, when the aqueous concentration of HCHO is relatively high) and  $O_3$  (L74). The reactions of  $HO_2+O_2^-$  (L7) and  $OH_2+O_2$  (L5) have positive impacts on S(IV), because they compete with L105 and L75 by consuming  $HO_2$  and  $H_2O_2$ , thus lower the conversion rates of S(IV) to S(VI) via L105 and L75. Effect of  $CH_2(OH)_2+OH$  (L50) and  $O_3+O_2^-$  (L13) on S(IV) and S(VI) can be either positive or negative. L50 consumes OH and produces  $HO_2$ , reducing the rates of aqueous oxidations of S(IV) involving OH but increasing those involving  $HO_2$  and  $O_2^-$ . Under the Remote conditions, L13 produces  $OH_2^-$ , which increases the pH and reduces the rates of some pH-dependent reactions that contribute to S(IV) conversion such as L7. Thus L13 has a positive effect on aqueous-phase S(IV). Being a major source for aqueous OH under the Marine conditions (and various polluted conditions, see below), on the other hand, the reaction of  $O_3+O_2^-$  (L13) increases the rates of L79 and L105, thus accelerates the conversion of S(IV) to S(VI).

Under various polluted conditions, the reactions of NO+HO<sub>2</sub> (G28) and OH+NO<sub>2</sub> (G26) become the dominant sinks for R<sub>x</sub>O<sub>y</sub>. Gas-phase H<sub>2</sub>O<sub>2</sub> dramatically decreases due to relatively high NO<sub>x</sub> and the rapid scavenging by cloud droplets. Due to the decreased gaseous H<sub>2</sub>O<sub>2</sub> concentration and its high reactivity in the aqueous-phase solution, the aqueous-phase  $H_2O_2$  concentration reaches its maximum ( (3-7) x 10 -5 M) within the first minute then substantially drops to ~10 -7 M within 10-40 minutes. As a result, the aqueousphase reaction of HSO<sub>3</sub>-+H<sub>2</sub>O<sub>2</sub> (L75) is no longer a dominant reaction for conversion of S(IV) to S(VI) after about 40 minutes. Unlike Remote and Marine atmospheres in which L75 acts as an important sink for aqueous-phase OH via depletion of H2O2 and dominates the conversion of S(IV) to S(VI) throughout the simulation, L75 becomes a positively-influential reaction for both aqueous-phase OH and S(IV) by decreasing pH and affecting pH-dependent reactions when aqueous H<sub>2</sub>O<sub>2</sub> concentrations are substantially lowered (i.e., after ~40 minutes) under the polluted conditions. Instead, S(IV)+HO<sub>2</sub> (L105) becomes the primary pathway for S(IV) conversion. Other important conversion reactions include oxidations of S(IV) by O<sub>2</sub> (catalyzed by Fe<sup>3+</sup> and Mn<sup>2+</sup>) (L77); oxidations of HSO<sub>3</sub>- by SO<sub>5</sub>- (L80), SO<sub>4</sub>- (L91), OH (L79), NO<sub>3</sub> (L108), and NO<sub>2</sub> (L109); as well as oxidations of SO<sub>3</sub> <sup>2-</sup> by SO<sub>5</sub>- (L81), HCHO (L112) and O<sub>3</sub> (L74). Other positively-influential reactions are  $SO_5^- + O_2^-$  (L82),  $CH_2(OH)_2 + OH$  (L50) and  $HO_2 + O_2^-$  (L7). These reactions can lower the reaction rates of L105 and L79 through reducing dissolved OH and HO2, thus indirectly reduce the conversion rates of S(IV).

While these results are generally consistent with the work of Pandis and Seinfeld (1989), notable differences are the direct impact of  $HO_2$  and  $H_2O_2$  on S(IV) and S(VI) through directly converting S(IV) to S(VI) (L105 and L75) and the indirect impact of  $HO_2$  on S(IV) by lowering the dissolved  $HO_2$  (=  $HO_2+O_2$ -) (L7). The reaction of S(IV) with  $HO_2$  (L105) is the second important pathway for S(IV) conversion under the relatively clean conditions and becomes the first important one under various polluted conditions, whereas it has little effect on S(IV) in the work of Pandis and Seinfeld (1989). L75 consistently decreases the aqueous-phase S(IV) in the work of Pandis and Seinfeld (1989), whereas it can affect S(IV) in a positive way at low  $H_2O_2$  concentrations (<  $10^{-7}$  M) under our modeled polluted conditions. This is because the predicted aqueous-phase  $HO_2$  concentrations under the five modeled conditions range from (0.9-4) x  $10^{-8}$  molecules/cm<sup>3</sup>, which are much higher than a value of ~1.0 x  $10^{-9}$  predicted by Pandis and Seinfeld (1989).

In addition, during the first 20 minutes the predicted aqueous  $H_2O_2$  concentrations significantly decrease from 7.0 x  $10^{-5}$  to 2.0 x  $10^{-5}$  M, from 5 x  $10^{-5}$  to 1.7 x  $10^{-6}$  M, and from 2.9 x  $10^{-5}$  to 2.7x  $10^{-8}$  M under the Rural, Urban and Heavily-polluted conditions, respectively. In contrast, the predicted aqueous  $H_2O_2$  concentrations in Pandis and Seinfeld (1989) maintained at a level of (0.9-5) x  $10^{-5}$  throughout the 2-hr simulation. The difference in predicted aqueous-phase  $HO_2$  and  $H_2O_2$  concentrations causes a relatively high conversion rate of S(IV) to S(VI) via L105 and sometimes an opposite effect of L75 under our modeled conditions. In addition to these differences, our simulation predicts a positive impact of  $HO_2+O_2$ - (L7) on S(IV), which is just the opposite to that of Pandis and Seinfeld (1989). The negative effect is because this reaction produces  $H_2O_2$ , thus accelerates the oxidation of S(IV) to S(VI) via L75, as interpreted by Pandis and Seinfeld (1989). In our cases, its negative effect on S(IV) through increasing  $H_2O_2$  is overridden by its positive effect through decreasing  $HO_2$  (via L105). Therefore, the net impact of the  $HO_2+O_2$ - reaction is a higher S(IV) and a lower S(VI).

## 4.2.2.2 Sensitivity to Gas-Aqueous Mass Transfer Coefficients

**Equilibrium Constants.** Table 12 shows the sensitivities of the predicted gas-phase O<sub>3</sub> concentrations with respect to Henry's Law constants and dissociation equilibrium constants under various conditions. The dissolution of HO<sub>2</sub> (E10) is the dominant negatively-influential reaction on O<sub>3</sub> under all conditions except for the Heavily-polluted conditions under which the gas-phase HO<sub>2</sub> has the lowest average concentrations of 2 x 10<sup>7</sup> molecules cm<sup>-3</sup> (i.e., 8 x 10 <sup>-4</sup> ppb). Other negatively-influential dissolution equilibria include those of O<sub>3</sub> (E17), H<sub>2</sub>O<sub>2</sub> (E11), HCHO (E12), and OH (E18). The Henry's law constants of NO<sub>2</sub> (E13) and HCOOH (E5) have positive effects on gaseous O<sub>3</sub> because these two equilibria tend to increase (rather than decrease) gas-phase NO<sub>2</sub> and HCOOH under cloudy conditions. Henry's law constants of HCHO (E12), NO<sub>2</sub>(E13), HCOOH (E5) and SO<sub>2</sub> (E3) become increasingly important as their gas-phase concentrations become higher. The dissociation of HO<sub>2</sub> (D2) plays a dominant role under all modeled conditions, and the importance of the dissociation of H<sub>2</sub>SO<sub>3</sub> (D10 and D11), HCHO (D9) and HCOOH (D16) increases as their aqueous-phase concentrations increase.

Mass Accommodation Coefficients. The rates of heterogeneous uptake and chemical transformation of species in clouds are determined by the overall mass transfer processes consisting of gasphase diffusion, interfacial mass accommodation, effective aqueous-phase solution and reaction, and aqueous-phase diffusion of dissolved species away from the particle surface (Schwartz, 1986; Worsnop et al., 1989). The mass transfer of a specific species can be controlled by any of the above processes, depending on the relative importance of these processes. Under our modeled conditions, most species do not experience appreciable mass-transfer limitation. Some oxidants such as OH, HO<sub>2</sub> and NO<sub>3</sub> are limited by the interfacial mass transfer process due to high solubility and/or high aqueous-phase effective first-order reaction rates. However, the effects of changing individual mass accommodation coefficients,  $\alpha_i$ , on overall predictions are mostly negligible. Sensitivity analyses show that most gaseous species except for NO<sub>3</sub>, N<sub>2</sub>O<sub>5</sub>, HNO<sub>3</sub>, H<sub>2</sub>O<sub>2</sub> and CH<sub>3</sub>OOH are not sensitive to changes in  $\alpha_i$ , with sensitivities less than 1.0 x 10<sup>-4</sup>. This is in consistence

with the findings of other researchers (Schwartz, 1986; Pandis and Seinfeld, 1989). The partitioning of CH<sub>3</sub>OOH, H<sub>2</sub>O<sub>2</sub> and N<sub>2</sub>O<sub>5</sub> between gas and aqueous-phase is appreciably affected by their own  $\alpha_i$ , recognizing that the uncertainty factors in the measured  $\alpha_i$  vary from 2-3 (DeMore et al., 1994). Their concentrations can decrease by 5%, 9% and 35% when doubling their own  $\alpha_i$ , respectively. The effects of  $\alpha_i$  on aqueous species are just the opposite to those on correspondingly gases. For example, as an result of higher  $\alpha$  of OH, HO<sub>2</sub>, HNO<sub>2</sub>, the aqueous-phase OH, HO<sub>2</sub>, H<sub>2</sub>O<sub>2</sub>, C<sub>2</sub>O<sub>3</sub>, CH<sub>3</sub>O<sub>2</sub>, and CH<sub>3</sub>COOH increase due to accelerated mass transfer from the gas to the aqueous-phase. None of these changes, however, has an appreciable influence on gaseous PAN, O<sub>3</sub> and indicators involving NO<sub>y</sub> and O<sub>3</sub> such as NO<sub>y</sub> and O<sub>3</sub>/NO<sub>z</sub>. It is noted that, however, changes in  $\alpha_i$  can appreciably affect indicators that involve H<sub>2</sub>O<sub>2</sub>, HNO<sub>3</sub> and HCHO, such as H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub>, H<sub>2</sub>O<sub>2</sub>/HCHO, H<sub>2</sub>O<sub>2</sub>/NO<sub>y</sub>, HCHO/HNO<sub>3</sub>, HCHO/NO<sub>y</sub> and O<sub>3</sub>/HNO<sub>3</sub>, with sensitivities on the order of 1.0 x 10-2. For example, doubling  $\alpha_{\text{HNO3}}$  under all polluted conditions can increase H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub> by ~8.8%. This is because both H<sub>2</sub>O<sub>2</sub> and HNO<sub>3</sub> are relatively sensitive to  $\alpha_i$ .

## 4.2.2.3 Sensitivity to Other Model Parameters

Sensitivities of major gaseous species and indicators to other model parameters such as temperature, R.H., cloud droplet size and liquid water ( $V_c$ ) are shown in Table 13. Temperature is the predominant parameter affecting overall model predictions under all conditions. A higher temperature causes increases in  $SO_2$ ,  $HNO_3$ ,  $H_2SO_4$ , OH,  $HO_2$ , PAN,  $O_3$ , total nitrate ( $TNO_3$ ) and decreases in total sulfate ( $TSO_4^2$ -). While an increase in temperature tends to increase the gas and aqueous-phase source and sink reaction rates of precursors, it decreases their solubilities in the aqueous-phase, thus reduces the rate of mass transfer from the gas-phase to aqueous-phase. Gas-phase concentrations of most species increase as a result of a net increase in their gaseous production rates and/or a decrease in their solubilities in the aqueous-phase. The lower  $TSO_4^{2-}$  is due to an increase in the sink of its precursors in the gas and/or aqueous-phase.

Either  $V_c$  or R.H. could be the second important parameter, depending on species and their dominant pathways. For species whose gas-phase reactions are dominant such as  $NO_x$ , PAN, ISOP, and OH, the effect of R.H. is more important. A higher R.H. produces higher water vapor and OH thus lower  $NO_x$  and ISOP. For species undergoing significant aqueous-phase reactions such as  $SO_2$ , HCHO, HO<sub>2</sub>, H<sub>2</sub>O<sub>2</sub>, O<sub>3</sub> and species strongly depending on aqueous-phase chemistry such as  $TSO_4^{2-}$  and  $TNO_3^-$ ,  $V_c$  is found to be the second important parameter. An increase in  $V_c$  greatly increases the gas to aqueous-phase mass transfer and the pH of the solution, thus affects many important species. As a result of a higher  $V_c$ , aqueous-phase HNO<sub>3</sub>, PAN,  $SO_2$ ,  $H_2SO_4$ , OH,  $HO_2$ , HCHO and  $O_3$  rapidly increase, lowering their corresponding gas-phase concentrations. Although the total amount of sulfate and nitrate also increases, their total concentrations ( $TNO_3^-$  and  $TSO_4^{2-}$ ) decrease due to higher dilution. In addition, a higher  $V_c$  results in higher  $NO_x$  under most conditions via lowering their oxidation rates by OH,  $HO_2$ , and  $O_3$ . These results are consistent with those of Pandis and Seinfeld (1989).

The cloud droplet size has an effect comparable to those of  $V_c$  and R.H. for most species under the Remote, Marine and Rural conditions, it becomes the least important parameter under the Urban and Heavily-polluted conditions. A larger droplet radius has an opposite effect to that of a higher  $V_c$ , i.e.,

decreasing the gas to aqueous-phase mass transfer rate. As a result, there remains higher gas-phase concentrations for most species under most conditions. However, gas-phase concentrations of some species such as  $NO_x$ , PAN,  $SO_2$ , OH,  $HO_2$ , ISOP and  $O_3$  can actually decrease under certain conditions. This is because there are net increases in the rates of their sinks despite of the decreasing mass transfer rate from the gas to aqueous-phase.

The impact of solar radiation,  $H_2O$  or temperature on indicator correlations has not been explored (Sillman, 1995). In our study, various indicators are also found to be highly sensitive to changes in temperature and moderately sensitive to changes in  $V_c$ , R.H., and cloud droplet size. An increase in temperature or R.H. tends to increase  $H_2O_2/HNO_3$ , but decreases both  $NO_y$  and  $O_3/NO_z$  under all polluted conditions. While an increase in  $V_c$  always increases these indicators, cloud droplet size affects them in a variety of ways, depending on the preexisting gas-phase concentrations. Comparison of Table 13 with Table 9 shows that some species and indicators show opposite sensitivities to temperature in the presence of clouds. For example, an increase in temperature results in a lower  $SO_2$  under all clear air conditions, but a higher  $SO_2$  under various cloudy conditions, implying a significant role of sulfur aqueous-phase chemistry on gas-phase  $SO_2$ . Temperature has a positive impact on  $NO_y$  and a negative effect on  $H_2O_2/HNO_3$  under clear air conditions. However, its effect becomes the opposite under cloudy conditions. Further study is needed to fully understand the role of temperature in the mixed-phase chemistry.

## 4.2.3 Aerosol Surface Chemistry

#### 4.2.3.1 Sensitivity to Individual Species Uptake Coefficients

The overall heterogeneous loss rates on aerosols strongly depend on individual uptake coefficients,  $\gamma_i$ . Unlike the sensitivities to  $\alpha_i$  under cloudy conditions, the predicted sensitivities of various species concentrations to  $\gamma_i$  in the presence of aerosols are higher by 1-3 orders of magnitude. For those species that are assumed to be directly scavenged to the aerosol surface, their concentrations are always anti-correlated with their own  $\gamma_i$  because higher  $\gamma_i$  deplete more gases onto the surface. For example, an increase in  $\gamma_{O3}$  always decreases gas-phase O<sub>3</sub>. Reactive species are also found to be affected by surface uptake of other species. The most influential aerosol surface reactions to overall model predictions include uptake of HCHO, O<sub>3</sub>, HO<sub>2</sub>, HNO<sub>2</sub>, HNO<sub>3</sub>, NO, NO<sub>2</sub>, NO<sub>3</sub>, N<sub>2</sub>O<sub>5</sub>, PAN, OH, H<sub>2</sub>O<sub>2</sub>, SO<sub>2</sub>, CH<sub>3</sub>O<sub>2</sub>, CH<sub>3</sub>OOH and CH<sub>3</sub>COOH.

Table 14 shows the sensitivities of various species including reactive nitrogen, odd hydrogen and reactive hydrocarbon to  $\gamma_i$  of the most influential species (with  $\overline{S}_{kl} > 1.0 \times 10^{-4}$ ) under the Heavily-polluted aerosol conditions. For reactive nitrogen species, the most influential  $\gamma_i$  are those of HCHO, O<sub>3</sub>, NO<sub>2</sub>, and N<sub>2</sub>O<sub>5</sub> in addition to their own  $\gamma_i$ . An increase of any of these  $\gamma_i$  either increases or decreases predicted gaseous species concentrations depending on its effects on the formation and destruction rates of these species. For example, an increase in  $\gamma_{\text{HCHO}}$  or  $\gamma_{\text{O3}}$  can reduce all reactive nitrogen species except for NO. This can be interpreted by the dominant reactions of these species. The dominant sinks for NO are O<sub>3</sub>+NO (G3), C<sub>2</sub>O<sub>3</sub>+NO (G46) and HO<sub>2</sub>+NO (G28) under this condition. An increase in  $\gamma_i$  of O<sub>3</sub> and HCHO can

cause lower gaseous  $O_3$  and HCHO, respectively. Since photolysis of HCHO (G38) is the dominant source of HO<sub>2</sub>, less HCHO results in less HO<sub>2</sub> in the gas-phase. As the result of lower  $O_3$  and HO<sub>2</sub>, the rates of G3 and G28 are lower, which results in a higher NO. On the contrary, G3, G28 and G38 act as sources for NO<sub>2</sub>, NO<sub>3</sub>, and N<sub>2</sub>O<sub>5</sub>. Lower gaseous O<sub>3</sub> and HCHO always cause lower formation rates of these species. For HNO<sub>3</sub> and PAN, a lower HCHO reduces the rate of G38, a dominant positively-influential reaction on both, thus leads to lower HNO<sub>3</sub> and PAN production. A lower O<sub>3</sub> causes lower OH and C<sub>2</sub>O<sub>3</sub> radicals, thus causes lower HNO<sub>3</sub> and PAN production through decreasing the rates of their sources OH+NO<sub>2</sub> (G26) and C<sub>2</sub>O<sub>3</sub>+NO<sub>2</sub> (G47), respectively. Other species that have appreciable impacts on predicted NO<sub>y</sub> species concentrations when changing their  $\gamma_i$  include NO, HNO<sub>3</sub>, HO<sub>2</sub>, and PAN.

For odd hydrogen species, the most important species are HCHO,  $O_3$ , HNO<sub>2</sub>, NO, NO<sub>2</sub>, HO<sub>2</sub>, H<sub>2</sub>O<sub>2</sub> and CH<sub>3</sub>OOH.  $\gamma_i$  of NO, NO<sub>2</sub>, N<sub>2</sub>O<sub>5</sub>, HNO<sub>3</sub> and SO<sub>2</sub> always have positive effects on their concentrations because the reactions of NO<sub>x</sub> and SO<sub>2</sub> with R<sub>x</sub>O<sub>y</sub> (e.g., G22, G26, G28, G29, G47, G115) are major sinks for OH. Higher uptake rates for NO<sub>x</sub> and SO<sub>2</sub> result in lower rates of these sinks thus higher R<sub>x</sub>O<sub>y</sub> species. Uptake of N<sub>2</sub>O<sub>5</sub> and HNO<sub>3</sub> on aerosol provides additional sinks for NO<sub>x</sub> and indirectly affects R<sub>x</sub>O<sub>y</sub> species. Lowering the rates of sources (e.g., G9-11, G38, G23, G32, G33, G48, and G118), on the other hand, O<sub>3</sub>, HCHO, HNO<sub>2</sub>, HO<sub>2</sub>, PAN, and CH<sub>3</sub>O<sub>2</sub> always have negative effects on R<sub>x</sub>O<sub>y</sub> species.

The ROG species assumed to be directly taken by aerosol surfaces include HCHO, HCOOH, CH<sub>3</sub>COOH, CH<sub>3</sub>O<sub>2</sub>, CH<sub>3</sub>OH, CH<sub>3</sub>OOH, and CH<sub>3</sub>COOOH. Concentrations of all ROG species are found to be not only directly affected by uptake of these species but also indirectly affected by uptake of other species such as HNO<sub>2</sub>, HO<sub>2</sub>, O<sub>3</sub>, NO, NO<sub>2</sub>, SO<sub>2</sub>, H<sub>2</sub>O<sub>2</sub>, and PAN. Increases in the  $\gamma_i$  of HCHO, HNO<sub>2</sub>, HO<sub>2</sub>, O<sub>3</sub>, and H<sub>2</sub>O<sub>2</sub> always decrease HCHO, ALD<sub>2</sub>, CRES, OPEN, HCOOH, CH<sub>3</sub>COOH and CH<sub>3</sub>OH but increase concentrations of other reactive ROG such as PAR, OLE, ETH, TOL, XYL and ISOP. Among all influential species, HCHO is the predominant one, with its effects 1-2 orders of magnitude higher than others. The effect of the lower gaseous HCHO due to a faster direct surface uptake on other species depends on their predominant reactions. For PAR, OLE, ETH, TOL, XYL and ISOP, their reactions with OH are the predominant pathway. In addition, photolysis of HCHO (G38) has a negative effect on these species, more HCHO uptake reduces rate of G38 thus lowers OH and HO<sub>2</sub>, resulting in a lower rate of sink via OH thus higher hydrocarbons in the gas-phase. For ALD<sub>2</sub>, CRES, OPEN, HCOOH, CH<sub>3</sub>COOH and CH<sub>3</sub>OH, G38 has a positive influence. While the lower rate of G38 results in the lower HO<sub>2</sub> and OH, which tends to increase their concentrations through reduction of their depletion rates by OH (e.g., G43, G66, G70), it also reduces the formation rates of these species such as CH<sub>3</sub>COOH through reducing organic radicals. In addition, lower OH and HO<sub>2</sub> also decrease the formation rate of ALD<sub>2</sub> through decreasing the rates of hydrocarbon interconversion reactions such as OLE, PAR and ETH converted to ALD<sub>2</sub> through G57, G58, G52, and G61. As a net result of this competition, the latter effect is predominant, i.e., lowering ALD<sub>2</sub>, CRES, OPEN, HCOOH, CH<sub>3</sub>COOH and CH<sub>3</sub>OH when increasing  $\gamma_{HCHO}$ .

Table 15(a) shows the sensitivity of  $O_3$  to  $\gamma_i$  of the most influential species under various aerosol conditions. The predicted  $O_3$  becomes more sensitive to changes in  $\gamma_i$  when the pre-existing atmospheric conditions become more polluted (i.e., more aerosol surface available). For example, doubling  $\gamma_{HCHO}$  and

 $\gamma_{O3}$  has almost no effect on O<sub>3</sub> under the Remote conditions, but it decreases O<sub>3</sub> by 6.3% and 10.4% under Although the mechanism remains unclear, the direct the Heavily-polluted conditions, respectively. destruction of O<sub>3</sub> on aerosols is found to be a dominant surface reaction under all modeled conditions. Stephens et al. (1986) found that the uptake of O<sub>3</sub> most likely occurs on carbonaceous or soot particles on which the adsorbed oxygen atom, which is produced from collision of O<sub>3</sub> with carbon aerosols, can combine with another adsorbed oxygen atom to form O<sub>2</sub>. In addition, one or two oxygen atoms may combine with a carbon atom to produce CO or CO<sub>2</sub>. They further determined that  $\gamma_{O3}$  on carbonaceous surfaces may be initially high but decreases more than a factor of 10 after repeated exposure resulting in an average  $\gamma_{O3}$  of 7 x 10-4, due probably to a saturation of surface sites by oxygenated groups. Fendel et al. (1995) also found that submicron carbon or iron aerosols can destroy  $O_3$  efficiently, with a  $\gamma_{O3}$  of 4 x 10-4. In this study, a  $\gamma_{O3}$  of 1 x 10<sup>-4</sup> was used, which is the lower limit in laboratory measurements. The predicted decreases in  $O_3$ concentrations due to 29 aerosol surface uptake reactions are 3%, 12.4% and 26.6% under the Rural, Urban and Heavily-polluted conditions, respectively, of which 18.2%, 37.0% and 39.9% are found to be caused by the direct O<sub>3</sub> destruction on aerosols, respectively. The rest of O<sub>3</sub> decrease is caused by the decrease in O<sub>3</sub> precursors such as NO<sub>2</sub>, HO<sub>2</sub> and HCHO. An increase in  $\gamma_i$  of these species also decreases O<sub>3</sub> because these surface uptake can significantly reduce the rates of O<sub>3</sub>-producing reactions (e.g., G1, G28, G38, G57, G61, and G69). On the contrary, higher  $\gamma_{NO}$  and  $\gamma_{SO2}$  tend to slightly increase O<sub>3</sub> because their surface reactions compete with their corresponding gas-phase reactions and decrease the rates of the gaseous reactions that contribute to O<sub>3</sub> destruction (e.g., G3, G22, and G115). As a result of all aerosol surface uptake, the O<sub>3</sub> formation rate decreases from 2.2 to 1.6 ppb hr<sup>-1</sup>, from 9.07 to 4.0 ppb hr<sup>-1</sup>, from 5.88 to 3.55 ppb hr<sup>-1</sup> under the Rural, Urban and Heavily-polluted conditions, respectively, as compared to the corresponding clear air conditions.

Since some reactive  $NO_v$  species such as  $N_2O_5$  and  $HNO_3$  and  $R_xO_v$  species such as  $H_2O_2$  and CH<sub>3</sub>OOH are highly sensitive to changes in  $\gamma_i$ , the indicators involving these species are expected to respond to these changes. While NO<sub>v</sub> is much less sensitive with most sensitivities less than 1.0 x 10-3, the sensitivities of O<sub>3</sub>/NO<sub>2</sub> and H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub> are 1-2 orders of magnitude higher, as shown in Table 15 (b) through (d). O<sub>3</sub>/NO<sub>z</sub> is sensitive to uptake of HNO<sub>3</sub>, HO<sub>2</sub>, H<sub>2</sub>O<sub>2</sub>, HCHO, NO<sub>2</sub>, PAN and O<sub>3</sub>. Similar to the effect on O<sub>3</sub>, an increase in γ<sub>O3</sub> always causes a decrease in O<sub>3</sub>/NO<sub>2</sub>. Higher uptake rates of NO<sub>2</sub> and N<sub>2</sub>O<sub>5</sub> can either increase (i.e., Rural and Urban) or decrease (i.e., Heavily-polluted) O<sub>3</sub>/NO<sub>2</sub>, depending on their combined effects on O<sub>3</sub> and NO<sub>2</sub>. The impacts of HCHO, HNO<sub>2</sub>, HO<sub>2</sub>, HO<sub>2</sub>, CH<sub>3</sub>O<sub>2</sub>, NO, and SO<sub>2</sub> on O<sub>3</sub>/NO<sub>z</sub>, however, are just the opposite as compared to those on O<sub>3</sub>. In addition, O<sub>3</sub>/NO<sub>z</sub> becomes sensitive to changes in  $\gamma_{HNO3}$  and  $\gamma_{PAN}$ , which have little effect on  $O_3$  formation. Since G38 is the dominant positively-influential reaction for both NO<sub>z</sub> and O<sub>3</sub>, a decrease in gaseous HCHO concentration reduces the photolytic rate of HCHO, thus decreases NO<sub>z</sub> and O<sub>3</sub>. Similarly, decreases in H<sub>2</sub>O<sub>2</sub>, CH<sub>3</sub>O<sub>2</sub> and HO<sub>2</sub> tend to decrease OH and O<sub>3</sub> concentrations. But they also decrease NO<sub>z</sub> concentrations by reducing the formation rates of NO<sub>3</sub>, N<sub>2</sub>O<sub>5</sub>, HNO<sub>3</sub> and PAN through reactions with OH. Therefore, the net effect of decreases in HCHO, HO<sub>2</sub>, H<sub>2</sub>O<sub>2</sub> and CH<sub>3</sub>O<sub>2</sub> is an increase in O<sub>3</sub>/NO<sub>z</sub>. The responses of H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub> are even more significant because both  $H_2O_2$  and  $HNO_3$  are highly sensitive to changes in their own  $\gamma_i$ . While an increase

in  $\gamma_{H2O2}$  greatly decreases gaseous  $H_2O_2$  thus reduces the indicator, an increase in  $\gamma_{HNO3}$  decreases gaseous  $HNO_3$  thus causes an increase in the indicator. In addition, this indicator is also sensitive to uptake of HCHO,  $HO_2$ ,  $O_3$ ,  $NO_x$ ,  $N_2O_5$ ,  $HNO_2$ , and  $SO_2$ .

## 4.2.3.2 Sensitivity to Temperature and Relative Humidity

Sensitivities of most gaseous species to temperature and R.H. under various aerosol conditions are similar to those under the corresponding clear air conditions shown in Table 9, with slightly higher values. Compared to clear air conditions, however, significant changes in both the magnitudes and the signs are found in sensitivities of  $NO_y$  and  $H_2O_2/HNO_3$ . In particular, a higher temperature causes a decrease in  $NO_y$  under all polluted conditions and an increase in  $H_2O_2/HNO_3$  under the Heavily-polluted conditions. A higher R.H. causes a higher  $H_2O_2/HNO_3$  under the Urban and Heavily-polluted conditions. These are just the opposite to those under the clear air conditions. These changes are similar to those under cloudy conditions (see Table 13), indicating that heterogeneous uptake of  $H_2O_2$  and  $NO_y$  species are important to their sensitivities, especially under conditions with large aerosol surfaces.

## 4.2.4 Effect of Heterogeneous Chemistry on Gas-Phase Chemistry

## 4.2.4.1 Effect of Clouds and Aerosols on O<sub>3</sub>-Precursor Relations

Gas-Phase Reaction Rate Constants. Heterogeneous chemistry alters the sensitivities of O<sub>3</sub> to various gas-phase reactions through changing their amplitudes and the signs. Figure 4 shows the comparison of the sensitivities of O<sub>3</sub> to gas-phase reaction rate constants in the absence and presence of clouds and aerosols under the Remote and Heavily-polluted conditions after a 2-hr simulation. Under the Remote conditions, cloud chemistry is the dominant process and causes significant changes in sensitivities to gasphase reactions, whereas the changes in sensitivities are mainly caused by aerosol surface reactions under the Heavily-polluted conditions. The sensitivities are generally smaller under the cloudy and aerosol conditions than those under clear air conditions, rationalizing a lower total gas-phase oxidizing capacity due to additional sinks for many O<sub>3</sub> precursors provided by clouds and aerosols. For example, the sensitivities of reactions of DMS, DMSO and CH<sub>3</sub>SO<sub>2</sub>H with OH (G82, G89, and G97, respectively) radicals are much lower under the cloudy and aerosol conditions because of lower OH levels. The effect of the gas-phase reaction CO+OH (G36) on O<sub>3</sub> formation has not only a different magnitude but also an opposite sign under the Remote cloudy conditions relative to the clear air conditions. It becomes negative under the cloudy conditions. This is because that HO<sub>2</sub> produced by G36 is not converted to OH via G28 under cloudy conditions, instead, it dissolves into cloud droplets, which increases the aqueous-phase concentration of  $O_{2}$ -, thus speeds up the important aqueous-phase sink of  $O_3$  through its reactions with  $O_2$ - (L13).

 $O_3$ -Precursor Relations. The presence of clouds and aerosols not only changes concentrations of  $O_3$  and its precursors, but also their relations provided that heterogeneous chemistry plays a significant role in

determining their mass budget. Figure 5 shows the comparison of the sensitivities of O<sub>3</sub> to species initial concentrations with and without clouds and aerosols under various polluted conditions after 2-hr simulation. O<sub>3</sub> increases with an increase in NO<sub>2</sub> in the presence of both clouds and aerosols under all conditions and in the presence of each alone under some conditions, which is just the opposite to that in clear air conditions. Under clear air conditions, the formation of HNO<sub>3</sub> (G26) and PAN (G47) through oxidation of NO<sub>2</sub> by OH and C<sub>2</sub>O<sub>3</sub> are predominant sink reactions for NO<sub>2</sub>. An increase in NO<sub>2</sub> results in a lower OH, thus a lower O<sub>3</sub>. Although OH still decreases with increasing NO<sub>2</sub> (due to R26) in the presence of clouds and/or aerosols, the fate of OH is no longer the dominant factor of determining the effect of NO<sub>2</sub> on O<sub>3</sub> production. The heterogeneous uptake and subsequent aqueous-phase reactions that affect NO2 and the relative importance of its gas-phase sinks via G26, G47, and G1 determine its correlation with O3. When there are sufficient surface areas, OH and C<sub>2</sub>O<sub>3</sub> are reduced due to lower levels of their precursors such as H<sub>2</sub>O<sub>2</sub>, HCHO and ALD<sub>2</sub> and the direct surface uptake of OH. This reduction could be so significant (20-48%, see Table 5) under some conditions (i.e., under the Rural and Urban cloudy condition, under the Urban and Heavilypolluted aerosol conditions, and under all polluted conditions with presence of both cloud and aerosols) that G26 and G47 are no longer the major sinks for NO<sub>2</sub>. Instead, G1 becomes the overwhelming sink for NO<sub>2</sub> and leads to O<sub>3</sub> formation followed by G2. Thus, O<sub>3</sub> formation increases with an increase in initial NO<sub>2</sub> under most cloudy and/or pre-existing aerosol conditions. In addition to the change in O<sub>3</sub>-NO<sub>2</sub> relations, O<sub>3</sub> becomes slightly more sensitive to NO under the Heavily-polluted conditions and less sensitive to initial O<sub>3</sub> and various ROG species such HCHO and ALD<sub>2</sub> under all conditions. For example, for a doubled HCHO, the predicted O<sub>3</sub> only increases by 15.4%, 1.8% and 1.9% in air with clouds alone, aerosols alone and both clouds and aerosols, respectively, as compared to 21.1% in clear air. Higher sensitivity of O<sub>3</sub> to initial NO is due to a slightly higher sensitivity to the rate constant of NO+O<sub>3</sub> (G3). O<sub>3</sub> sensitivities to various ROG are lower because the TOC is reduced and O3 sensitivities to most ROG reactions such as G38 and G43 are appreciably lower, as shown in Figure 4.

## 4.2.4.2 Effect of Clouds and Aerosols on Sensitivity of Indicators

Significant changes are found in sensitivities of indicators to both reaction rate constants and species initial concentrations due to the cloud chemistry and aerosol surface uptake. Although the sensitivities of NO<sub>y</sub> to most gas-phase reactions rate constants are greatly increased in the presence of clouds and/or aerosols, their magnitudes are still smaller by 1-3 orders than those of O<sub>3</sub>/NO<sub>z</sub> and H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub>, indicating that NO<sub>y</sub> is relatively insensitive to reaction rate constants. Figure 6 compares the sensitivities of O<sub>3</sub>/NO<sub>z</sub> and H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub> to gas-phase reaction rate constants with and without clouds and aerosols under the Heavily-polluted conditions. O<sub>3</sub>/NO<sub>z</sub> becomes more sensitive to G9-G11, G43-G47, G58, G69, G74, and G76, and less sensitive to G1, G3, G38, G39, G52, and G57. The most significant change is that the impact of the rate constant of NO<sub>2</sub>+OH (G26) on O<sub>3</sub>/NO<sub>z</sub> changes from negative to positive. Under clear air conditions, HNO<sub>3</sub> produced via this reaction remains in the gas-phase and largely increases the denominator, NO<sub>z</sub>. Thus a higher rate constant of G26 actually decreases the indicator. When there are sufficient cloud

droplets and/or aerosol particles,  $HNO_3$  can be quickly taken onto these surfaces as soon as it is produced in the gas-phase. In the meanwhile, other  $NO_z$  species such as  $NO_3$ ,  $N_2O_5$  and PAN can undergo similar uptake and subsequent aqueous-phase reactions, therefore decrease gas-phase  $NO_z$ . This decrease is larger than the simultaneous decrease in  $O_3$  due to the surface uptake. As a result, the indicator increases with a higher rate constant of G26. The sensitivities of  $H_2O_2/HNO_3$  to many reaction rate constants in the presence of clouds and aerosols are significantly changed in both magnitudes and signs because both  $H_2O_2$  and  $HNO_3$  can be effectively scavenged by clouds and aerosols, and these surface uptake and aqueous-phase chemistry become the predominant pathways for  $H_2O_2$  and  $HNO_3$ .

The changes in these sensitivities cause substantial changes in the sensitivities of the indicators to initial species concentrations, as shown in Figure 7. NO<sub>v</sub> becomes less sensitive to NO<sub>2</sub>, but more sensitive to initial O<sub>3</sub>, NO, and ROG species such as HCHO, ALD<sub>2</sub>, and OLE. O<sub>3</sub>/NO<sub>z</sub> becomes much more sensitive to changes in NO<sub>2</sub>, ALD<sub>2</sub>, OLE, TOL, XYL and ISOP and less sensitive to changes in O<sub>3</sub>, NO, HCHO, and PAR. The changes in sensitivities of H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub> to various species are even more significant. H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub> becomes extremely sensitive (with  $S_{kl} > 0.9$ ) to initial NO, NO<sub>2</sub>, O<sub>3</sub>, and OLE and almost insensitive to H<sub>2</sub>O<sub>2</sub>. In addition, the impact of initial O<sub>3</sub>, HCHO, OLE, ETH, TOL, XYL and ISOP on H<sub>2</sub>O<sub>2</sub>/HNO<sub>3</sub> changes from negative under clear air conditions to positive under cloud and/or aerosol conditions. Cloud chemistry and aerosol surface uptake cause significant changes in dominant reactions of H<sub>2</sub>O<sub>2</sub>. H<sub>2</sub>O<sub>2</sub> is no longer a reservoir for OH and HO<sub>2</sub> (due to G34 and G35). Once produced, H<sub>2</sub>O<sub>2</sub> can be rapidly scavenged into cloud droplets or onto aerosol surfaces. In addition, other odd hydrogen species such as OH and HO<sub>2</sub> can also be effectively removed by clouds and aerosols and undergo rapid aqueous-phase reactions. As a result, the gaseous H<sub>2</sub>O<sub>2</sub> and OH and HO<sub>2</sub> are substantially lower. While higher NO<sub>2</sub> increases HNO<sub>3</sub> via G26, it also causes further decreases in OH (due to G26), thus a lower H<sub>2</sub>O<sub>2</sub> and a subsequent significant decrease in the indicator. The extremely high sensitivity to initial NO, NO<sub>2</sub>, and OLE was caused by the much higher sensitivities to their corresponding reactions (i.e., G26, G28, and G57) shown in Figure 6(b). Similarly, the opposite effect of initial O<sub>3</sub>, HCHO, OLE, ETH, TOL, XYL and ISOP is due to the corresponding changes in sensitivities to their dominant reactions (i.e., G9-11, G38, G57, G58, G61, G63, G69 and G76).

#### 5. Conclusions

A mixed-phase chemistry box model consisting of detailed gas- and aqueous-phase chemistry and a simplified treatment of aerosol surface chemistry was used to study heterogeneous chemistry and its effect on gas-phase chemistry under a variety of atmospheric conditions. A subsequent comprehensive sensitivity analysis was conducted to evaluate the sensitivity of overall model predictions and identify the most influential model parameters that affect O<sub>3</sub> formation, O<sub>3</sub>-precursor relations and photochemical indicators in the mixed-phase chemical system. The results show that heterogeneous processes associated with clouds and aerosols not only reduce many gas-phase species concentrations and the total oxidizing capacity but alter O<sub>3</sub>-precursor relations. Decreases in O<sub>3</sub> and PAN concentrations can be up to 27% and 38%, respectively, with

up to 100% decrease in their formation rates under typical atmospheric conditions. The magnitude of the heterogeneous effects is very sensitive to temperature, R.H., cloud water content and individual uptake coefficients. The effects of clouds and aerosols can be accelerated by a higher water content and higher individual uptake coefficients, respectively. Temperature and R.H. can influence both cloud and aerosol processes in a variety of ways.

The most influential gas-phase reactions to O<sub>3</sub> formation under the relatively clean conditions include the photolytic reactions of NO<sub>2</sub>, O<sub>3</sub>, HONO, H<sub>2</sub>O<sub>2</sub>, and HCHO; the conversion of NO to NO<sub>2</sub> by HO<sub>2</sub>, O<sub>3</sub>, CH<sub>3</sub>O<sub>2</sub>, CH<sub>3</sub>SCH<sub>2</sub>OO and C<sub>2</sub>O<sub>3</sub>; the generation and inter-conversion of OH, HO<sub>2</sub> and RO<sub>2</sub> via reactions O(<sup>1</sup>D)+H<sub>2</sub>O, O<sub>3</sub>+HO<sub>2</sub>, and reactions of OH with CH<sub>4</sub>, O<sub>3</sub>, CO, DMSO, DMS, CH<sub>3</sub>OOH, HCHO, ALD<sub>2</sub>, OLE, ETH and PAR; and the formation and/or dissociation of oxidants and acids such as CH<sub>3</sub>OOH, HNO<sub>3</sub>, H<sub>2</sub>O<sub>2</sub>, HNO<sub>4</sub>, CH<sub>3</sub>SO<sub>2</sub>, H<sub>2</sub>SO<sub>4</sub>, HONO, PAN, and CH<sub>3</sub>SCH<sub>2</sub>OO. The most influential gas-phase reactions under various polluted conditions are the photolytic reactions of NO<sub>2</sub>, O<sub>3</sub>, HONO, H<sub>2</sub>O<sub>2</sub>, HCHO, ALD<sub>2</sub>, OPEN, and MGLY; the conversion of NO to NO<sub>2</sub> by HO<sub>2</sub>, O<sub>3</sub>, C<sub>2</sub>O<sub>3</sub> and TO<sub>2</sub>; the generation and interconversion of OH, HO<sub>2</sub> and RO<sub>2</sub> through reactions O(<sup>1</sup>D)+H<sub>2</sub>O, OLE+O<sub>3</sub>, and reactions of OH with CO, CH<sub>4</sub>, ETH, ALD<sub>2</sub>, PAR, OLE, ISOP, TOL, HCHO, XYL, and MGLY; and the formation and dissociation of HNO<sub>3</sub>, H<sub>2</sub>SO<sub>4</sub>, HNO<sub>4</sub>, H<sub>2</sub>O<sub>2</sub> and PAN.

Cloud chemistry is a dominant heterogeneous process in relatively clean and moderately polluted atmospheres. The most influential aqueous-phase reactions to O<sub>3</sub> formation under the relatively clean conditions are oxygen-hydrogen reactions of O<sub>2</sub>- with O<sub>3</sub> and HO<sub>2</sub>, HO<sub>2</sub>+HO<sub>2</sub>, H<sub>2</sub>O<sub>2</sub>+O<sub>3</sub>, OH+H<sub>2</sub>O<sub>2</sub>, OH+HO<sub>2</sub>, photolysis of H<sub>2</sub>O<sub>2</sub>; sulfur reactions HSO<sub>3</sub>-+H<sub>2</sub>O<sub>2</sub>, SO<sub>3</sub><sup>2</sup>-+O<sub>3</sub>, S(IV)+HO<sub>2</sub>, HSO<sub>3</sub><sup>-</sup>+O<sub>3</sub>, HOCH<sub>2</sub>SO<sub>3</sub>-+OH; methane oxidation CH<sub>2</sub>(OH)+OH and CH<sub>3</sub>O<sub>2</sub>+O<sub>2</sub>-; carbonate reactions of CO<sub>3</sub>- with H<sub>2</sub>O<sub>2</sub> and O<sub>2</sub>- and HCO<sub>3</sub>+O<sub>2</sub>-; and nitrate reactions NO<sub>3</sub>-+Cl<sup>-</sup>. The most influential aqueous-phase reactions under various polluted conditions include oxygen-hydrogen reactions of O<sub>2</sub>- with O<sub>3</sub> and HO<sub>2</sub>, OH+HO<sub>2</sub>, HO<sub>2</sub>+HO<sub>2</sub>; sulfur reactions of HSO<sub>3</sub>- with H<sub>2</sub>O<sub>2</sub>, OH, SO<sub>5</sub>-, NO<sub>2</sub>, and O<sub>3</sub>, reactions of SO<sub>3</sub><sup>2</sup>- with O<sub>3</sub>, SO<sub>5</sub>- and HCHO, reactions of S(IV) with HO<sub>2</sub> and O<sub>2</sub> (catalyzed by Mn<sup>2+</sup> and Fe<sup>3+</sup>), reactions of SO<sub>5</sub>- with SO<sub>5</sub>- and O<sub>2</sub>-, HOCH<sub>2</sub>SO<sub>3</sub>-+OH, and SO<sub>4</sub>-+HCOO-; nitrate reactions NO<sub>3</sub>+HO<sub>2</sub>, NO<sub>2</sub>+NO<sub>2</sub> and photolysis of NO<sub>3</sub>-; methane oxidation CH<sub>2</sub>(OH)+OH and HCOO-+OH; carbonate reaction HCO<sub>3</sub>-+O<sub>2</sub>-. Among these, the reactions of O<sub>2</sub>- with dissolved HO<sub>2</sub> and O<sub>3</sub> are mainly responsible for the decrease in O<sub>3</sub> formation under most cloudy conditions.

Aerosols can be more effective surfaces than clouds in polluted areas when the total surface area is larger than  $1000~\mu\text{m}^2~\text{cm}^{-3}$ . The aerosol surface chemistry is very sensitive to the individual uptake coefficients, which are subject to large uncertainties. Among the 29 condensing species, the heterogeneous uptake of O<sub>3</sub>, HCHO, HO<sub>2</sub>, NO<sub>2</sub>, NO, N<sub>2</sub>O<sub>5</sub>, HNO<sub>2</sub>, HNO<sub>3</sub>, H<sub>2</sub>O<sub>2</sub>, SO<sub>2</sub>, CH<sub>3</sub>O<sub>2</sub>, PAN, CH<sub>3</sub>OOH and CH<sub>3</sub>COOH are the most influential aerosol surface reactions. In particular, the significant decrease in O<sub>3</sub> formation in the presence of aerosols is mainly caused by the heterogeneous uptake of O<sub>3</sub>, HCHO, HO<sub>2</sub>, H<sub>2</sub>O<sub>2</sub> and NO<sub>2</sub>. Further laboratory and field studies are urgently needed to precisely quantify this important parameter and quantify the rates and product yields of these heterogeneous reactions on aerosols.

The  $O_3$  formation is also sensitive to a number of mass transfer coefficients and physical parameters. Under the relatively clean conditions, the Henry's law constants of  $HO_2$ ,  $O_3$ ,  $H_2O_2$ , and HCHO; the

dissociation equilibrium constants of HO<sub>2</sub> and H<sub>2</sub>SO<sub>3</sub>; and the uptake coefficients of O<sub>3</sub>, H<sub>2</sub>O<sub>2</sub>, and HO<sub>2</sub> are the most important parameters in determining the mass transfer between the gas and the condensed-phases. Under various polluted conditions, the most influential parameters include the Henry's law constants of HO<sub>2</sub>, HCHO, O<sub>3</sub>, SO<sub>2</sub>, NO<sub>2</sub>, and HCOOH; the dissociation equilibrium constants of HO<sub>2</sub>, H<sub>2</sub>SO<sub>3</sub>, HCHO, and HCOOH; and the uptake coefficients of O<sub>3</sub>, HCHO, HO<sub>2</sub>, NO<sub>2</sub>, NO, N<sub>2</sub>O<sub>5</sub>, HNO<sub>2</sub>, HNO<sub>3</sub>, PAN, H<sub>2</sub>O<sub>2</sub>, CH<sub>3</sub>OOH, CH<sub>3</sub>COOH and SO<sub>2</sub>. Temperature, R.H., cloud water content and droplet size can largely affect O<sub>3</sub> and its precursors in a variety of ways. O<sub>3</sub> formation is relatively insensitive to changes in mass accommodation coefficients of individual species under all modeled conditions.

As a result of changes in gas-phase concentrations and the total oxidizing capacities, the magnitude of photochemical indicators and their sensitivity to model parameters are also largely affected. Presence of clouds and/or aerosols can increase  $O_3/NO_z$  and  $O_3/HNO_3$ , decrease  $NO_y$ ,  $H_2O_2/NO_y$ ,  $HCHO/NO_y$  ( $HNO_3/H_2O_2/(NO_x/ROG)$ ), and affect  $O_3/NO_y$  and  $H_2O_2/HNO_3$  in both ways under all modeled conditions. The indicators  $NO_y$  and  $O_3/NO_y$  are relatively insensitive to most model parameters, whereas indicators involving  $H_2O_2$ ,  $HNO_3$ , HCHO and  $NO_z$  are highly sensitive to changes in initial species concentrations, reaction rate constants, equilibrium constants, uptake coefficients, temperature, R.H., cloud droplet size and water content. While their most influential reactions are similar to those of  $O_3$ , their relative impacts may be different both in signs and amplitudes. Indicators generally have higher sensitivities to ROG reactions and lower sensitivities to  $NO_x$  reactions under polluted conditions than less polluted conditions, which coincides with the nature of the indicators as good markers for the  $O_3$ - $NO_x$ -ROG sensitivity. In addition, sensitivities of the indicators to model parameters can be significantly changed when including cloud and aerosol chemistry. Caution should be taken when applying the established threshold values of indicators to determine  $NO_x$ - or ROG-sensitive  $O_3$  in regions where large cloud coverage and high aerosol loading are frequently observed.

Our results indicate that the effect of aerosols is similar and comparable to that of clouds when the aerosol loading is large (e.g., in urban atmospheres). However, the predicted aerosol effect is based on the assumption of a non-reversible mass transfer between gas and aerosol surfaces, which may not always be the case in the atmosphere. Future work is needed to refine aerosol treatments by accounting for gas-aerosol equilibria, saturation and reactions on various aerosols. Our results also demonstrate a need to further study the role of heterogeneous chemistry in tropospheric chemistry through sensitivity analysis of a 3-D model. In particular, their impacts on O<sub>3</sub> formation, O<sub>3</sub>-precursor relations and photochemical indicators need to be further studied in 3-D models. Furthermore, we have only evaluated model responses to changes in the chemical mechanism, and there are many other uncertain parameters in the "real" atmosphere such as emission and meteorological conditions. Results from 3-D sensitivity studies will be extremely useful in evaluating O<sub>3</sub>-NO<sub>x</sub>-ROG sensitivities and developing integrated control strategies for O<sub>3</sub> and particulate matters in non-attainment regions.

Our work show that large-scale sensitivity analysis of complex computer models is feasible with automatic differentiation tools, although the fully automatic approach currently is quite compute-intensive. Automatic differentiation is a technology in its infancy, so further improvements in tools are expected.

However, exploitation of high-level program or algorithmic structure has been shown to have a dramatic impact on computational cost in certain application studies. Thus, as more experience is gained in the use of AD tools, we expect that synergy between improvements in AD tools and high-level user understanding will decrease the computational cost by orders of magnitude. With improved aerosol treatment and AD tools, a comprehensive 3-D sensitivity analysis will be possible and greatly enhance our understanding of mixed-phase chemical system and their impact on photochemistry in the "real" atmosphere.

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Table 1. Chemical species included in the MaTChM.

Notice of the content of the conte	N.T.	0 1 .	D	N.	A 1 1 .
2         Nitrogan tiroxide (tritate radical)         NO2         2         HNO2           4         Daintrogen pentoxide         N205         4         HEXOCH           8         Nitrogan (control decidence)         N205         4         HEXOCH           8         Nitros acid         HONO         5         HOCOH           9         Notes acid         HONO         7         COC           9         Oxygen attorn (singlet)         O(1D)         8         MSA           11         Ozoge         O3         11         H200           11         Hydroxy Tadical         OH         10         H02           12         Hydroxy Tadical         HD2         12         HCH           13         Hydroxy Tadical         HD2         12         HCH           14         Hydroxy Tadical         HCH         HCH         NO           15         Carbon monoxide         CO         15         NO           16         Hydroxy Tadical (CHSCO)OOT)         C203         17         O3           17         Peroxysey Tadical (CHSCO)OOT)         C203         17         O3           18         Peroxysey Tadical (CHSCO)OOT)         C203         17 </td <td>No.</td> <td>Gas-phase species Nitric oxide</td> <td>Representation NO</td> <td>No 1</td> <td>Aqueous and aerosol species HNO3</td>	No.	Gas-phase species Nitric oxide	Representation NO	No 1	Aqueous and aerosol species HNO3
1   100					
Matrice   Matr					
Mixica acid (HOZNOL2)			N2O5	4	H2SO4
Perxy partic acid (HOZNOZ)					
80         Oxygen atom (singlet)         OLDD         8         MSA           10         Hydroxyl radical         OH         10         HO2           12         Hydroxyl radical         II OO         11         H2OC           12         Hydroper persoxide         H2OC         13         NO2           14         Formatichysic (CH2-O)         HCHO         11         NO3           14         Formatichysic (CH2-O)         HCHO         15         NO3           15         High molecular weight alchydes (RCHO, R>H)         LDZ         16         PAN           16         High molecular weight alchydes (RCHO, R>H)         LDZ         16         PAN           17         Persaysogal radical CH2GCOOON2         PAN         18         OH           18         Persoyagal radical CH2GCOON2         PAR         20         CH3OH           19         Paraffic action book (C-C)         PAR         20         CH3OH           21         Scondary organic oxy radical         ROR         21         DMS           22         NO4-actinate operation         NO22         22         DMSO           23         Olefinic carbon bond (C-C)         QL         EBAR         20         CH3OH <td></td> <td></td> <td></td> <td></td> <td></td>					
90         Oxygen arom (triplet)         Oxygen arom (triplet)         H   10   100   1					
10			. ,		
11   10   10   10   11   12   12   13   14   14   15   15   15   15   15   15			` '		
13   Hydrogen peroxide   H2O2   15   NO2					
14   Formaldehyde (CH2-O)					
15 Carbon monoxide         CO         15 No3           16 High molecular weight aldehydes (RCHO, R>H)         AlD2         16 PAN           17 Peroxyacyl radical (CHSCO)OOP)         C203         17 O3           18 Peroxyacyl radical (CHSCO)OON)2         PAN         18 OH           19 NO40-NO2 operation         XO2         19 CH302           20 Paraffin carbon bond (C-C)         PAR         20 CH30H           21 Scoondary organic oxy radical         ROR         21 DMS           22 NO40-oittrate operation         XO2N         22 DMSO           23 Olefinic carbon bond (C-C)         OLE         23 DMSO2           24 Ethene(CHZ-CH2)         ETH         24 CH30OH           25 Toluene(C6H5-CH3)         TOL         25 CH3C(O)OOH           26 Cresol and higher molecular weight phenols         CRES         26 NZOS           27 Toluene-hydroxyr radical adduct         TO2         27 NTR           28 Methylphenoxy radical         CROS         28 HC1           29 High molecular weight aromatic oxidation ring fragment         OPEN         29 CL           31 Methylphoxal (CH3CC)C(O)H)         ROB         20 CL           32 Methylphenoxy radical         CH30C         28 PCL           33 Methylphenoxy radical         CH3SCQ         29 CO3					
16         High molecular weight aldehydes (RCHO, R>H)         ALD2         16         PAN           17         Peroxyacyl intract (CHSCIO)OOY)         PAN         18         OH           18         Peroxyacyl intract (CHSCIO)OON(2)         PAN         18         OH           20         PARTIIN carbon bond (C-C)         PAR         20         CH3OH           21         Secondary organic oxy radical         ROR         21         DMS           22         NO-to-nitrate operation         XOZN         22         DMSO           21         Secondary organic oxy radical         CRE         23         DMSO           24         Ethene(CH2-CH2)         ETH         24         CH3OOH           25         Toleane-hydroxyl radical adduct         CRES         26         NZOS           26         Cresol and higher molecular weight phenols         CRES         28         HCI           27         Toleane-hydroxyl radical adduct         CRO         28         HCI           28         Heigh molecular weight phenols         CRES         26         NZOS           29         High molecular weight phenols         RRE         CRO         28         HCI           20         High molecular weight adduct <td></td> <td></td> <td></td> <td></td> <td></td>					
Persoyacy  radical (CH3CO)OOO)					
18		• •			
19   NO-to-NO2 operation	18		PAN	18	OH
21         Secondary organic oxy radical         ROR         21         DMS           22         NOt-on-intrate operation         XOZN         22         DMSO           23         Olefinic carbon bond (C=C)         OLE         23         DMSO           24         Ethen (CH2-CH2)         TOL         25         CH3CO)OOH           25         Toluene (CH5-CH3)         TOL         25         CH3CO)OOH           26         Creso and higher molecular weight phenols         CRES         26         NCH2CH0Honoxy radical           27         Toluene-hydroxyl radical adduct         CRO         28         HCI           29         High molecular weight anomatic oxidation ring fragment         OPEN         29         CI           30         Nylen (CH3-CH3CO)C(O)H)         MGLY         SV         SV         CH3CGH4-CH32)         AU         CH3CGH4-CH32)         CO3*           31         unknown organic oxidation product of nitrogen species         NTR         No         Ionic species           33         unknown organic oxidation product of nitrogen species         NTR         No         Ionic species           34         Dimethyl sulfonic (CH3C)SO3         MSA         4         SOS*         OCO3*         SOS*           <					
22         No-to-nitrate operation         XOZN         22         DMSOC           24         Eithene(CH2=CH2)         ETH         24         CH3000H           25         Tolune Carbon bond (G=C)         ETH         24         CH3000H           26         Cresol and higher molecular weight phenols         CRES         26         N205           27         Tolune-hydroxyl radical adduct         TO2         27         NTR           28         Methylphenoxy radical         CRO         28         HCI           30         Sylene(C6H4-(CH3)2)         XYL         Well           31         Methylgycval (CH3C(O)C(O)H)         MGLY         SOP           32         Isoprene         ISOP         ISOP           33         usknown organic oxidation product of nitrogen species         NTR         NO         Ionia: species           34         Dimethyl sulfoxil (c(H3)2S         DMS         1         CO3*         CO3*           35         Methyl sulfoxyl acid         CH3SO2         2         HOCH2SO3*           36         Methyl sulfoxyl acid (CH3)2SO         DMSO2         6         CH2(OH)2           40         Methyl sulfoxide (CH3)2SO         DMSO2         6         CH2(OH)2		Paraffin carbon bond (C-C)			
23         Olefinic carbon bond (C=C)         OLE         23         DMSO2           44         Ethen (CH2-CH2)         ETH         24         CH30OH           25         Toluene (CH3-CH3)         TOL         25         CH3C(O)OH           27         Toluene-hydroxyl radical adduct         TO2         27         NTR           28         Methylphenoxy radical         CRO         28         HC1           29         High molecular weight aromatic oxidation ring fragment         OPEN         29         Cl           30         Xylen (CG4H4-CCH32)         XYL         XYL         V           21         Isopree         ISOP         ISOP         V           31         unknown organic oxidation product of nitrogen species         NTR         N         Ionic species           33         unknown organic oxidation product of nitrogen species         NTR         N         Ionic species           34         Dimethyl sulfonic acid (CH3C303H)         MSA         4         SOG*           35         Methyl sulfonic acid (CH3SO3H)         MSA         4         SOG*           36         Methyl sulfoxide (CH3)2SO         DMSO         5         CH2COH)2           40         Methyl sulfoxide (CH3)2SO					
24 Ibhene(CH2=CH2)         ETH Land (CH3 CH3)         24 CH3 COOH         CH3 COOH         25 CH3C(O)OOH         26 CH3C(O)OOH         26 CH3C(O)OOH         26 CH3C(O)OOH         27 CH3 CH3C(O)OOH         27 CH3C(CH3 CH3)         27 CH3C(CH3 CH3)         28 CH3C(O)OOH         28 CH3C(O)OOH         28 CH3C(O)OOH         28 CH3C(CH3 CH3 CH3)         28 CH3C(CH3 CH3)         28 CH3C(CH3 CH3)         29 CH3C(CH3 CH3)         28 CH3C(CH3 CH3)         28 CH3C(CH3 CH3)         28 CH3C(CH3 CH3)         29 CH3C(CH3 CH3)         28 CH3C(CH3 CH3)         29 CH3C(CH3 CH3)         29 CH3C(CH3 CH3)         29 CH3C(CH3 CH3)         20 CH3C(CH3)		1			
25         Crisoland higher molecular weight phenols         CRES         25         CH3C(O)OGH           27         Toluene-hydroxyl radical adduct         TO2         27         NTR           28         Methylehnoxy radical         CRO         28         HC1           29         High molecular weight aromatic oxidation ring fragment         OPEN         29         CI           30         Xylenet(CH4-(CH3)2)         MCHylglyoxal (CH3C(O)C(O)H)         MGLY         MGLY           31         Inknown organic oxidation product of nitrogen species         NTR         No         lonic species           34         Dimethyl sulfford, (CH3)2S         DMS         1         CO3*           35         Methyl sulfonic radical         CH3SO3         3         SO4*           36         Methyl sulfoxide (CH3)2SO         DMSO         5         HSO5*           37         Methane sulfonic acid (CH3SO3H)         MSA         4         SO5*           38         Dimethyl sulfoxide (CH3)2SO         DMSO2         6         CH2(OH1)2           40         Methyl sulfoxyl acid         CH3SO(2H         7         H*           41         Methyl sulfoxyl acid         CH3SO(2H         7         H*           42         Dim					
26         Cresol and higher molecular weight phenols         CRES         26         N205           27         Totlane-hydroxyl radical adduct         CRO         28         HCI           28         High phenoxy radical         OPEN         29         CI           30         Xylenc(C6H4-(CH3)2)         XYL         XYL           31         Methyl ploxoal (CH3C(O)C(O)H)         MGLY         **           32         Isoprene         ISOP         1         C03*           33         unknown organic oxidation product of nitrogen species         NTM         N         Monte species           34         Dimethyl sulfonic oxidation product of nitrogen species         DMS         1         C03*           35         Methyl sulfonic radical         CH3SO2         2         HOCH2SO3*           37         Methyl sulfonic radical         CH3SO2         2         HCOCH2SO3*           38         Dimethyl sulfonic acid (CH3)2SO         DMSO         5         HSO5*           40         Methyl sulfonyl selfone (CH3)2SO         DMSO         6         CH2(OH)2*           41         Methyl sulfonyl selfone (CH3)2SO         MS         NH4*           42         Dimethyl sulfonic product of sulfur secies         CH3SO(DA)					
27         Toluene-hydroxyl radical adduct         TOQ         27         NTR           29         Methylphenoxy radical         CRO         28         HCI           29         High molecular weight aromatic oxidation ring fragment         OPEN         29         CI           30         Xylene CGH4-C(H3)2)         YXI         VXI           31         Incorect         ISOP         ISOP           32         unknown organic oxidation product of nitrogen species         NTR         No         Ionic species           33         unknown organic oxidation product of nitrogen species         NTR         No         Ionic species           34         Dimethyl sulfonyl sulforal         CH3SO2         2         HOCH2SO3*           36         Methyl sulfonic radical         CH3SO3         3         SO4*           37         Methane sulfonic acid (CH3SO3H)         MSA         4         SO5*           38         Dimethyl sulfonic acid (CH3SO2D)         DMSO         5         HSO5*           39         Dimethyl sulfonyl peroxyl radical         CH3SO2H         7         H*           40         Methyl sulfonyl peroxyl radical         CH3SO2DO         8         NH4*           41         Methyl sulfonyl peroxyl radical					` '
29 High molecular weight aromatic oxidation ring fragment         OPEN VIL (NGCH4-(CH3)2)         29 KI (NGCH4-(CH3)2)         CI (NGCH4-(CH3)2)         27 (NGCH4-(CH3)2)         27 (NGCH4-(CH3)2)         20 (NGCH4-(C	27	Toluene-hydroxyl radical adduct			
Nylene(CoH4-(CH3)2)		Methylphenoxy radical			
31 Intercepting (Amony)         Methylglyoxal (CH3C(O)C(O)H)         Methylglyoxal (CH3C(O)C(O)H)         Methyle (CH3)C(O)C(O)H)         Mode (Amony)         Intercepting (Amony)         Methyle (CH3)C(O)C(O)H)         Methyle (CH3)C(O)C(O)H)         Methyle (CH3)C(O)C(O)C(O)H)         Methyle (CH3)C(O)C(O)C(O)C(O)C(O)C(O)C(O)C(O)C(O)C(O		High molecular weight aromatic oxidation ring fragment		29	Cl
Sopreme   Sopr		• • • • • • • • • • • • • • • • • • • •			
33         unknown organic oxidation product of nitrogen species         NTR         No         lonic species           34         Dimethyl sulfide, (CH3)2S         DMS         1         CO3"           35         Methyl sulfonic radical         CH3SO2         2         HOCH2SO3"           36         Methyl sulfonic radical         CH3SO3         3         SO4"           37         Methane sulfonic acid (CH3)2SO1         DMSO         5         HSO5"           39         Dimethyl sulfonic (CH3)2SO2         DMSO2         6         CH2(OH)2           40         Methyl sulfonyl peroxyl radical         CH3SO(DOO         8         NH4"           41         Methyl sulfonyl peroxyl radical         CH3S(D)2OO         8         NH4"           42         Dimethyl sulfonyl peroxyl radical         CH3S(D)2CH2OO         10         Fe(OH)2"           43         Dimethyl sulfonyl peroxyl radical         CH3S(O)2CH2OO         10         Fe(OH)2"           44         Sulfuric acid         H2SO4         12         Mn2"           45         Sulfuric acid         H2SO4         12         Mn2"           46         Unkown oxidation product of sulfur species         SULF         13         OH           48					
35         Methyl sulfonic radical         CH3SO2         2         HOCH2SO3*           36         Methyl sulfonic radical         CH3SO3         3         SO4*           37         Methane sulfonic acid (CH3SO3H)         MSA         4         SO5*           38         Dimethyl sulfonic Acid (CH3)2SO2         DMSO2         6         CH2(OH)2           40         Methyl sulfonyl acid         CH3SO2H         7         H*           41         Methyl sulfonyl peroxyl radical         CH3S(O)2OO         8         NH4*           42         Dimethyl sulfonyl peroxyl radical         CH3S(O)2CH2OO         10         Fe(OH)2*           43         Dimethyl sulfonyl peroxyl radical         CH3S(O)2CH2OO         10         Fe(OH)2*           44         Sulfuri dioxide         SO2         11         Fe(OH)2*           45         Sulfuri dioxide         SO2         11         Fe(OH)2*           46         Unkown oxidation product of sulfur species         SULF         13         OH*           47         Methane         CH4         14         NO3*           48         Ethane         CH4         14         NO3*           49         Methyl peroxy radical formed from alkane         ETHP		1		No	Ionic species
35         Methyl sulfonic radical         CH3SO3         3         SO4*           36         Methyl sulfonic radical         CH3SO3         3         SO4*           37         Methane sulfonic acid (CH3SO3H)         MSA         4         SO5*           38         Dimethyl sulfonic (CH3)2SO2         DMSO2         6         CH2(OH)2           40         Methyl sulfonyl acid         CH3SO2H         7         H*           41         Methyl sulfonyl peroxyl radical         CH3S(O)2OO         8         NH4*           42         Dimethyl sulfonyl peroxyl radical         CH3S(O)2CH2OO         10         Fe(OH)2*           43         Dimethyl sulfonyl peroxyl radical         CH3S(O)2CH2OO         10         Fe(OH)2*           44         Sulfur dioxide         SO2         11         Fe(OH)2*           45         Sulfuric acid         H2SO4         12         Mn2*           46         Unkown oxidation product of sulfur species         SULF         13         OH*           47         Methane         CH4         14         NO3*           48         Ethane         CH3O         16         HSO3*           50         Ethyl peroxyl radical         CH3O         18         HSO4* <td>34</td> <td>Dimethyl sulfide, (CH3)2S</td> <td>DMS</td> <td>1</td> <td>*</td>	34	Dimethyl sulfide, (CH3)2S	DMS	1	*
36         Methyl sulfonic radical         CH3SO3         3         SO4*           37         Methane sulfonic acid (CH3SO3H)         MSA         4         SO5*           38         Dimethyl sulfoxide (CH3)2SOQ         DMSOQ         6         CH2(OH)2           40         Methyl sulfonyl c(CH3)2SOQ         DMSOQ         6         CH2(OH)2           40         Methyl sulfonyl peroxyl radical         CH3SCO2H         7         H*           41         Methyl sulfonyl peroxyl radical         CH3SCH2OO         9         Fe3**           42         Dimethyl sulfonyl peroxyl radical         CH3SCO)2CH2OO         10         Fe(OH)2**           43         Dimethyl sulfonyl peroxyl radical         CH3SCO)2CH2OO         10         Fe(OH)2**           44         Sulfuri doxide         SO2         11         Fe(OH)2**           45         Sulfuri doxide         SO2         11         Fe(OH)2**           46         Unkown oxidation product of sulfur species         SULF         13         OH*           47         Methane         CH4         14         NO3*           48         Ethane         CH3O         16         HSO3*           50         Ethyl peroxy radical formed from alkane         ETHP </td <td>35</td> <td>Methyl sulfonyl radical</td> <td>CH3SO2</td> <td>2</td> <td>HOCH2SO3</td>	35	Methyl sulfonyl radical	CH3SO2	2	HOCH2SO3
Methane sulfonic acid (CH3SO3H)   MSA	36	Methyl sulfonic radical	CH3SO3	3	
38         Dimethyl sulfoxide (CH3)2SO         DMSO         5         HSO5*           39         Dimethyl sulfone (CH3)2SO2         DMSO2         6         CH2(OH)2           40         Methyl sulfonyl acid         CH3SO2H         7         H*           41         Methyl sulfonyl peroxyl radical         CH3S(O)2OO         8         NH4*           42         Dimethyl sulfonyl peroxyl radical         CH3S(O)2CH2OO         10         Fe(OH)2*           43         Dimethyl sulfonyl peroxyl radical         CH3S(O)2CH2OO         10         Fe(OH)2*           44         Sulfur dioxide         SO2         11         Fe(OH)2*           45         Sulfur dioxide         SO2         11         Fe(OH)2*           46         Unkown oxidation product of sulfur species         SULF         13         OH           47         Methane         CH4         14         NO3*           48         Ethane         C2H6         15         NO2*           49         Methyl peroxyl radical         CH3O2         16         HSO3*           50         Ethyl peroxy radical formed from alkane         ETHP         17         SO3*           51         Methyl peroxide         CH3OOH         18 <td< td=""><td>37</td><td>Methane sulfonic acid (CH3SO3H)</td><td>MSA</td><td>4</td><td></td></td<>	37	Methane sulfonic acid (CH3SO3H)	MSA	4	
39         Dimethyl sulfone (CH3)2SO2         DMSO2 (CH3SO2H)         6 CH2(OH)2           40         Methyl sulfonyl acid         CH3SO2H         7 H <sup>+</sup> 41         Methyl sulfonyl peroxyl radical         CH3S(O)2OO         8 NH <sub>4</sub> <sup>+</sup> 42         Dimethyl sulfide peroxyl radical         CH3SCH2OO         9 Fe3 <sup>+</sup> 43         Dimethyl sulfonyl peroxyl radical         CH3S(O)2CH2OO         10 Fe(OH)2 <sup>+</sup> 44         Sulfur dioxide         SO2         11 Fe(OH)2 <sup>+</sup> 45         Sulfur dioxide         SO2         11 Fe(OH)2 <sup>+</sup> 46         Unkown oxidation product of sulfur species         SULF         13 OH*           47         Methane         CH4         14 NO3 <sup>-</sup> 48         Ethane         C2H6         15 NO2 <sup>-</sup> 49         Methyl peroxyl radical         CH3O2         16 HSO3 <sup>-</sup> 50         Ethyl peroxy radical formed from alkane         ETHP         17 SO3 <sup>=</sup> 51         Methyl peroxide         CH3OOH         18 HSO4 <sup>-</sup> 52         Acetic acid         CH3COOH         19 SO4 <sup>-</sup> 53         Formic acid         HCOOH         20 HCO3 <sup>-</sup> 54         Methanol         CH3OH	38		DMSO	5	
40         Methyl sulfonyl acid         CH3SO2H         7         H <sup>+</sup> 41         Methyl sulfonyl peroxyl radical         CH3S(O)2OO         8         NH4 <sup>+</sup> 42         Dimethyl sulfide peroxyl radical         CH3SCH2OO         9         Fe3 <sup>+</sup> 43         Dimethyl sulfonyl peroxyl radical         CH3S(O)2CH2OO         10         Fe(OH)2 <sup>+</sup> 44         Sulfur dioxide         SO2         11         Fe(OH)2 <sup>+</sup> 45         Sulfuric acid         H2SO4         12         Mn2 <sup>+</sup> 46         Unkown oxidation product of sulfur species         SULF         13         OH           47         Methane         CH4         14         NO3 <sup>-</sup> 48         Ethane         C2H6         15         NO2 <sup>-</sup> 49         Methyl peroxyl radical         CH3O2         16         HSO3 <sup>-</sup> 50         Ethyl peroxy radical formed from alkane         ETHP         17         SO3 <sup>-</sup> 51         Methyl peroxide         CH3OOH         18         HSO4 <sup>-</sup> 52         Acetic acid         CH3OOH         19         SO4 <sup>-</sup> 53         Formic acid         HCOOH         20         HCO3 <sup>-</sup> </td <td></td> <td>•</td> <td></td> <td></td> <td></td>		•			
Methyl sulfonyl peroxyl radical   CH3S(O)2OO   8					
42         Dimethyl sulfide peroxyl radical         CH3SCH2OO         9         Fe <sup>3+</sup> 43         Dimethyl sulfonyl peroxyl radical         CH3S(O)2CH2OO         10         Fe(OH) <sup>2+</sup> 44         Sulfur dioxide         SO2         11         Fe(OH) <sup>2+</sup> 45         Sulfuric acid         H2SO4         12         Mn <sup>2+</sup> 46         Unkown oxidation product of sulfur species         SULF         13         OHT           47         Methane         CH4         14         NO3 <sup>-</sup> 48         Ethane         C2H6         15         NO2 <sup>-</sup> 49         Methyl peroxyl radical         CH3O2         16         HSO3 <sup>-</sup> 50         Ethyl peroxyl radical formed from alkane         ETHP         17         SO3 <sup>-</sup> 51         Methyl peroxide         CH3OOH         18         HSO4 <sup>-</sup> 52         Acetic acid         CH3COOH         19         SO4 <sup>-</sup> 53         Formic acid         HCOOH         20         HCO3 <sup>-</sup> 54         Methanol         CH3OH         21         CO3 <sup>-</sup> 55         Carbon dioxide         CO2         22         O2 <sup>-</sup> 57		•			
43         Dimethyl sulfonyl peroxyl radical         CH3S(O)2CH2OO         10         Fe(OH) <sup>2+</sup> 44         Sulfur dioxide         SO2         11         Fe(OH) <sup>2+</sup> 45         Sulfuric acid         H2SO4         12         Mn <sup>2+</sup> 46         Unkown oxidation product of sulfur species         SULF         13         OH           47         Methane         CH4         14         NO3 <sup>-</sup> 48         Ethane         C2H6         15         NO2 <sup>-</sup> 49         Methyl peroxyl radical         CH3O2         16         HSO3 <sup>-</sup> 50         Ethyl peroxy radical formed from alkane         ETHP         17         SO3 <sup>=</sup> 51         Methyl peroxide         CH3OOH         18         HSO4 <sup>-</sup> 52         Acetic acid         CH3COOH         19         SO4 <sup>=</sup> 53         Formic acid         HCOOH         20         HCO3 <sup>-</sup> 54         Methanol         CH3OH         21         CO3 <sup>=</sup> 55         Carbon dioxide         CO2         22         O2 <sup>-</sup> 56         Ammonia         NH3         23         HO2 <sup>-</sup> 57         Hydrogen chloride </td <td>42</td> <td>Dimethyl sulfide peroxyl radical</td> <td>CH3SCH2OO</td> <td>9</td> <td>-</td>	42	Dimethyl sulfide peroxyl radical	CH3SCH2OO	9	-
44       Sulfur dioxide       SO2       11       Fe(OH)2+         45       Sulfuric acid       H2SO4       12       Mn2+         46       Unkown oxidation product of sulfur species       SULF       13       OH         47       Methane       CH4       14       NO3*         48       Ethane       C2H6       15       NO2*         49       Methyl peroxyl radical       CH3O2       16       HSO3*         50       Ethyl peroxy radical formed from alkane       ETHP       17       S03*=         51       Methyl peroxide       CH3OOH       18       HSO4*         52       Acetic acid       CH3COOH       19       SO4*=         53       Formic acid       HCOOH       20       HCO3*         54       Methanol       CH3OH       21       C03*=         55       Carbon dioxide       CO2       22       O2*         56       Ammonia       NH3       23       HO2*         57       Hydrogen chloride       HCI       24       HCOO*         No       Lumped gas-phase species       25       CH3COO*         1       Nitrogen oxides=NO+NO2       NOx       26       CH3SO3**					
H2SO4					
46					
47         Methane         CH4         14         NO3 <sup>-</sup> 48         Ethane         C2H6         15         NO2 <sup>-</sup> 49         Methyl peroxyl radical         CH3O2         16         HSO3 <sup>-</sup> 50         Ethyl peroxy radical formed from alkane         ETHP         17         SO3 <sup>-</sup> 51         Methyl peroxide         CH3OOH         18         HSO4 <sup>-</sup> 52         Acetic acid         CH3COOH         19         SO4 <sup>-</sup> 53         Formic acid         HCOOH         20         HCO3 <sup>-</sup> 54         Methanol         CH3OH         21         CO3 <sup>-</sup> 55         Carbon dioxide         CO2         22         O2 <sup>-</sup> 56         Ammonia         NH3         23         HO2 <sup>-</sup> 57         Hydrogen chloride         HCI         24         HCOO <sup>-</sup> NO         Lumped gas-phase species         25         CH3COO <sup>-</sup> 1         Nitrogen oxides=NO+NO2         NOx         26         CH3SO3 <sup>-</sup> 2         Total reactive nitrogen=NO+NO2+NO3+N2O5+HNO3+PAN+NTR         NOz         28         Cl <sup>-</sup> 3         Total reactive nitrogen=NO4-NO2+NO3+ROB F	45			12	$Mn^{2+}$
47       Methane       CH4       14       NO3 <sup>-</sup> 48       Ethane       C2H6       15       NO2 <sup>-</sup> 49       Methyl peroxyl radical       CH3O2       16       HSO3 <sup>-</sup> 50       Ethyl peroxy radical formed from alkane       ETHP       17       SO3 <sup>-</sup> 51       Methyl peroxide       CH3OOH       18       HSO4 <sup>-</sup> 52       Acetic acid       CH3COOH       19       SO4 <sup>-</sup> 53       Formic acid       HCOOH       20       HCO3 <sup>-</sup> 54       Methanol       CH3OH       21       CO3 <sup>-</sup> 55       Carbon dioxide       CO2       22       O2 <sup>-</sup> 56       Ammonia       NH3       23       HO2 <sup>-</sup> 57       Hydrogen chloride       HCI       24       HCOO <sup>-</sup> 1       Nitrogen oxides=NO+NO2       NOx       26       CH3SO3 <sup>-</sup> 2       Total reactive nitrogen=NO+NO2+NO3+N2O5+HNO3+PAN+NTR       NOz       28       CI <sup>-</sup> 3       Total non-NOx reactive nitrogen=NOy-NOx       NOz       28       CI <sup>-</sup>	46	Unkown oxidation product of sulfur species	SULF	13	OH-
48         Ethane         C2H6         15         NO2^-           49         Methyl peroxyl radical         CH3O2         16         HSO3^-           50         Ethyl peroxy radical formed from alkane         ETHP         17         SO3=           51         Methyl peroxide         CH3OOH         18         HSO4^-           52         Acetic acid         CH3COOH         19         SO4=           53         Formic acid         HCOOH         20         HCO3^-           54         Methanol         CH3OH         21         CO3=           55         Carbon dioxide         CO2         22         02^-           56         Ammonia         NH3         23         HO2^-           57         Hydrogen chloride         HCl         24         HCOO^-           NO         Lumped gas-phase species         25         CH3COO^-           1         Nitrogen oxides=NO+NO2         NOx         26         CH3SO3^-           2         Total reactive nitrogen=NO+NO2+NO3+N2O5+HNO3+PAN+NTR         NOz         28         CI^-           3         Total non-NOx reactive nitrogen=NOy-NOx         NOz         28         CI^-	47	Methane	CH4	14	NO3 <sup>-</sup>
49         Methyl peroxyl radical         CH3O2         16         HSO3 <sup>-</sup> 50         Ethyl peroxy radical formed from alkane         ETHP         17         SO3 <sup>-</sup> 51         Methyl peroxide         CH3OOH         18         HSO4 <sup>-</sup> 52         Acetic acid         CH3COOH         19         SO4 <sup>-</sup> 53         Formic acid         HCOOH         20         HCO3 <sup>-</sup> 54         Methanol         CH3OH         21         CO3 <sup>-</sup> 55         Carbon dioxide         CO2         22         O2 <sup>-</sup> 56         Ammonia         NH3         23         HO2 <sup>-</sup> 57         Hydrogen chloride         HCl         24         HCOO <sup>-</sup> No         Lumped gas-phase species         25         CH3COO <sup>-</sup> 1         Nitrogen oxides=NO+NO2         NOx         26         CH3SO3 <sup>-</sup> 2         Total reactive nitrogen=NO+NO2+NO3+N2O5+HNO3+PAN+NTR         NOz         28         CI <sup>-</sup> 3         Total non-NOx reactive nitrogen=NOy-NOx         NOz         28         CI <sup>-</sup>	48	Ethane	C2H6	15	
50         Ethyl peroxy radical formed from alkane         ETHP         17         SO3=           51         Methyl peroxide         CH3OOH         18         HSO4-           52         Acetic acid         CH3COOH         19         SO4=           53         Formic acid         HCOOH         20         HCO3-           54         Methanol         CH3OH         21         CO3=           55         Carbon dioxide         CO2         22         O2-           56         Ammonia         NH3         23         HO2-           57         Hydrogen chloride         HCI         24         HCOO-           NO         Lumped gas-phase species         25         CH3COO-           1         Nitrogen oxides=NO+NO2         NOx         26         CH3SO3-           2         Total reactive nitrogen=NO+NO2+NO3+N2O5+HNO3+PAN+NTR         NOy         27         CIOH-           3         Total non-NOx reactive nitrogen=NOy-NOx         NOz         28         CI-	49	Methyl peroxyl radical	CH3O2	16	
51         Methyl peroxide         CH3OOH         18         HSO4 <sup>-</sup> 52         Acetic acid         CH3COOH         19         SO4 <sup>=</sup> 53         Formic acid         HCOOH         20         HCO3 <sup>-</sup> 54         Methanol         CH3OH         21         CO3 <sup>=</sup> 55         Carbon dioxide         CO2         22         O2 <sup>-</sup> 56         Ammonia         NH3         23         HO2 <sup>-</sup> 57         Hydrogen chloride         HCl         24         HCOO <sup>-</sup> NO         Lumped gas-phase species         25         CH3COO <sup>-</sup> 1         Nitrogen oxides=NO+NO2         NOx         26         CH3SO3 <sup>-</sup> 2         Total reactive nitrogen=NO+NO2+NO3+N2O5+HNO3+PAN+NTR         NOy         27         CIOH <sup>-</sup> 3         Total non-NOx reactive nitrogen=NOy-NOx         NOz         28         Cl <sup>-</sup>					
52       Acetic acid       CH3COOH       19       SO4=         53       Formic acid       HCOOH       20       HCO3-         54       Methanol       CH3OH       21       CO3=         55       Carbon dioxide       CO2       22       O2-         56       Ammonia       NH3       23       HO2-         57       Hydrogen chloride       HCI       24       HCOO-         NO       Lumped gas-phase species       25       CH3COO-         1       Nitrogen oxides=NO+NO2       NOx       26       CH3SO3-         2       Total reactive nitrogen=NO+NO2+NO3+N2O5+HNO3+PAN+NTR       NOy       27       CIOH-         3       Total non-NOx reactive nitrogen=NOy-NOx       NOz       28       CI-					
53         Formic acid         HCOOH         20         HCO3 <sup>-</sup> 54         Methanol         CH3OH         21         CO3 <sup>=</sup> 55         Carbon dioxide         CO2         22         O2 <sup>-</sup> 56         Ammonia         NH3         23         HO2 <sup>-</sup> 57         Hydrogen chloride         HCI         24         HCOO <sup>-</sup> NO         Lumped gas-phase species         25         CH3COO <sup>-</sup> 1         Nitrogen oxides=NO+NO2         NOx         26         CH3SO3 <sup>-</sup> 2         Total reactive nitrogen=NO+NO2+NO3+N2O5+HNO3+PAN+NTR         NOy         27         CIOH <sup>-</sup> 3         Total non-NOx reactive nitrogen=NOy-NOx         NOz         28         Cl <sup>-</sup>		• •			
54         Methanol         CH3OH         21         CO3=           55         Carbon dioxide         CO2         22         O2-           56         Ammonia         NH3         23         HO2-           57         Hydrogen chloride         HCl         24         HCOO-           No         Lumped gas-phase species         25         CH3COO-           1         Nitrogen oxides=NO+NO2         NOx         26         CH3SO3-           2         Total reactive nitrogen=NO+NO2+NO3+N2O5+HNO3+PAN+NTR         NOy         27         CIOH-           3         Total non-NOx reactive nitrogen=NOy-NOx         NOz         28         Cl-					
55         Carbon dioxide         CO2         22         O2 <sup>-</sup> 56         Ammonia         NH3         23         HO2 <sup>-</sup> 57         Hydrogen chloride         HCl         24         HCOO <sup>-</sup> No         Lumped gas-phase species         25         CH3COO <sup>-</sup> 1         Nitrogen oxides=NO+NO2         NOx         26         CH3SO3 <sup>-</sup> 2         Total reactive nitrogen=NO+NO2+NO3+N2O5+HNO3+PAN+NTR         NOy         27         CIOH <sup>-</sup> 3         Total non-NOx reactive nitrogen=NOy-NOx         NOz         28         Cl <sup>-</sup>					HCO3 <sup>-</sup>
56         Ammonia         NH3         23         HO2 <sup>-</sup> 57         Hydrogen chloride         HCl         24         HCO0 <sup>-</sup> No         Lumped gas-phase species         25         CH3CO0 <sup>-</sup> 1         Nitrogen oxides=NO+NO2         NOx         26         CH3SO3 <sup>-</sup> 2         Total reactive nitrogen=NO+NO2+NO3+N2O5+HNO3+PAN+NTR         NOy         27         CIOH <sup>-</sup> 3         Total non-NOx reactive nitrogen=NOy-NOx         NOz         28         Cl <sup>-</sup>					CO3 <sup>=</sup>
57 Hydrogen chloride  No Lumped gas-phase species  1 Nitrogen oxides=NO+NO2  2 Total reactive nitrogen=NO+NO2+NO3+NO3+PAN+NTR  NOy  3 Total non-NOx reactive nitrogen=NOy-NOx  NOz  NOz  NOz  NOz  NOz  NOz  NOZ  NOZ	55	Carbon dioxide			O2 <sup>-</sup>
57     Hydrogen chloride     HCl     24     HCOO <sup>-</sup> No     Lumped gas-phase species     25     CH3COO <sup>-</sup> 1     Nitrogen oxides=NO+NO2     NOx     26     CH3SO3 <sup>-</sup> 2     Total reactive nitrogen=NO+NO2+NO3+N2O5+HNO3+PAN+NTR     NOy     27     ClOH <sup>-</sup> 3     Total non-NOx reactive nitrogen=NOy-NOx     NOz     28     Cl <sup>-</sup>	56	Ammonia	NH3	23	HO2 <sup>-</sup>
No Lumped gas-phase species  1 Nitrogen oxides=NO+NO2 NOx 26 CH3COO- 2 Total reactive nitrogen=NO+NO2+NO3+N2O5+HNO3+PAN+NTR NOy 27 CIOH- 3 Total non-NOx reactive nitrogen=NOy-NOx NOz 28 CI-	57	Hydrogen chloride	HC1	24	
Nitrogen oxides=NO+NO2  NOx  26 CH3SO3  Total reactive nitrogen=NO+NO2+NO3+N2O5+HNO3+PAN+NTR NOy  Total non-NOx reactive nitrogen=NOy-NOx NOz  NOz  NOz  NOz  NOz  NOz  NOz  N	No	Lumped gas-phase species		25	
2 Total reactive nitrogen=NO+NO2+NO3+N2O5+HNO3+PAN+NTR NOy 27 CIOH <sup>-</sup> 3 Total non-NOx reactive nitrogen=NOy-NOx NOz 28 CI <sup>-</sup>			NOx		
3 Total non-NOx reactive nitrogen=NOy-NOx NOz 28 Cl		-			
A Odd bydoner OH, HOA (MIZO) ETHIR (202) DOR (TO) (CRO		•	· ·		
4 Odd nydrogen=OH+HOZ+CH3OZ+E1HP+CZO3+KOK+TOZ+CKO KXOy 29 Cl2-		•			
	4	Oaa nyarogen=OH+HOZ+CH3OZ+ETHP+C2O3+ROR+TOZ+CRO	KxOy	29	Cl <sub>2</sub> -

Table 2. Mass accommodation coefficients ( $\alpha$ ) and uptake coefficients ( $\gamma$ ) of species in the MaTChM.

No	Species	α	Reference	$\gamma = \alpha_{min}$	Reference
1	HNO3	1.1E-1	Ponche et al., 1993	2.4E-3	Baldwin, 1982, see note [e]
2	HNO2	3.0E-2	Bongartz et al., 1994	5.0E-4	Kirchner et al., 1990, see note [d]
3	SO2	5.4E-2	Gardner et al., 1987	2.4E-4	Judeikis et al., 1978
4	H2SO4	2.0E-2	Dingenen and Raos, 1991	2.0E-4	see note [c]
5	НСООН	4.7E-2	Jayne et al., 1991	4.7E-4	see note [c]
6	СН3СООН	4.7E-2	see note [a]	4.7E-4	see note [a]
7	CO2	1.0E-4	see note [b]	1.0E-6	see note [c]
8	MSA	9.0E-2	DeBruyn et al., 1994	9.0E-4	see note [c]
9	NH3	9.7E-2	Ponche et al., 1993	9.7E-4	see note [c]
10	HO2	2.0E-1	Mozurkewich et al., 1987; Kurylo et al.,1994	1.0E-1	Hanson et al., 1992, see note [e]
11	H2O2	1.8E-1	Worsnop et al., 1989	7.8E-3	Baldwin, 1982, see note [e]
12	НСНО	2.0E-2	Jayne et al., 1992	1.0E-3	Jayne et al., 1992
13	NO2	6.3E-4	Lee and Tang, 1988	3.3E-5	Baldwin, 1982, see note [e]
14	NO	1.0E-5	Leu, 1988	1.0E-5	Baldwin, 1982, see note [e]
15	NO3	1.0E-2	see note [b]	1.0E-2	Thomas et al., 1989, see note [e]
16	PAN	1.0E-3	Kirchner et al., 1990	1.0E-4	Kirchner et al., 1990, see note [d]
17	O3	5.3E-4	Tang and Lee, 1987	1.0E-4	Fendel et al., 1995
18	OH	8.0E-2	Hanson et al., 1992	4.9E-3	Baldwin and Golden, 1980, see note [e]
19	CH3O2	5.0E-2	Dentener, 1993	5.0E-3	Dentener, 1993, see note [d]
20	СН3ОН	3.4E-2	Davidovits et al., 1993	3.4E-3	Davidovits et al., 1993, see note [d]
21	DMS	7.4E-2	see note [a]	7.4E-3	see note [a]
22	DMSO	7.4E-2	Davidovits et al., 1993	7.4E-3	Davidovits et al., 1993, see note [d]
23	DMSO2	7.9E-2	Davidovits et al., 1993	7.9E-3	Davidovits et al., 1993, see note [d]
24	СНЗООН	2.9E-2	Davidovits et al., 1993	2.9E-4	Davidovits et al., 1993, see note [c]
25	СНЗСОООН	2.9E-2	see note [a]	2.9E-4	see note [a]
26	N2O5	1.0E-1	Morzurkewich and Calvert, 1988	1.0E-1	Morzurkewich and Calvert, 1988
27	NTR	1.0E-6	Baldwin and Golden, 1980, see note [f]	1.0E-8	see note [c]
28	HCl	1.4E-1	Van Doren et al., 1990	1.0E-2	Kirchner et al., 1990
29	Cl	1.0E-4	Martin et al., 1980	3.0E-5	Martin et al., 1980

a αs and γs of DMS, CH3COOH and CH3COOH are assumed to be the same as those of DMSO, HCOOH and CH3OOH, respectively.

b For species with missing or insufficient  $\alpha$  and  $\gamma$  measurement data and for which measurements of analogous compounds do not exist, the values of  $\alpha$  are assumed to be 0.01 for soluble species, and 1.E-4 for less soluble species, respectively.

c  $\gamma$  is assumed to be a factor of 100 lower than the corresponding  $\alpha$  used in this work.

d  $\gamma$  is assumed to be a factor of 10 lower than the corresponding measured lower limit of  $\alpha$  reported in the reference.

e  $\gamma$  is assumed to be a factor of 10 higher than the corresponding measured lower limit of  $\gamma$  reported in the reference.

The mass accommodation coefficient of other nitrogen species (NTR) is assumed to be the same as that of atomic nitrogen in Baldwin and Golden, 1980.

Table 3. Heterogeneous uptake of various species on the surface of aerosols in MaTChM.

No	Heterogeneous surface uptake		Overall	heterogeneous loss rate,	$K_{pi}$ , s <sup>-1</sup>	
		Remote	Marine	Rural	Urban	Heavily-polluted
P1.	$HNO_3(g) \rightarrow HNO_3(p)$	2.46E-6	5.19E-6	2.51E-5	2.08E-4	5.90E-4
P2.	$HNO_2(g) \rightarrow HNO_2(p)$	5.39E-7	1.14E-6	5.30E-6	4.38E-5	1.24E-4
P3.	$SO_2(g) \rightarrow H_2SO_3(p)$	2.55E-7	5.39E-7	2.51E-6	2.07E-5	5.86E-5
P4.	$H_2SO_4(g) \rightarrow H_2SO_4(p)$	1.72E-7	3.63E-7	1.69E-6	1.39E-5	3.95E-5
P5.	$HCOOH(g) \rightarrow HCOOH(p)$	5.87E-7	1.24E-6	5.79E-6	4.78E-5	1.35E-4
P6.	$CH_3COOH(g) \rightarrow CH_3COOH(p)$	5.12E-7	1.08E-6	5.07E-6	4.19E-5	1.19E-4
P7.	$CO_2(g) \rightarrow H_2CO_3(p)$	1.29E-8	2.72E-8	1.26E-7	1.04E-6	2.95E-6
P8.	$MSA(g) \rightarrow MSA(p)$	7.71E-7	1.63E-6	7.66E-6	6.34E-5	1.79E-4
P9.	$NH_3(g) \rightarrow NH_4OH(p)$	1.92E-6	4.05E-6	1.95E-5	1.62E-4	4.59E-4
P10.	$HO_2(g) \to HO_2(p)$	5.33E-5	1.16E-4	1.20E-3	1.08E-2	3.25E-2
P11.	$H_2O_2(g) \to H_2O_2(p)$	9.28E-6	1.97E-5	1.08E-4	9.12E-4	2.60E-3
P12.	$HCHO(g) \rightarrow HCHO(p)$	1.50E-6	3.17E-6	1.52E-5	1.26E-4	3.56E-4
P13.	$NO_2(g) \rightarrow NO_2(p)$	4.16E-8	8.77E-8	4.07E-7	3.36E-6	9.51E-6
P14.	$NO(g) \rightarrow NO(p)$	2.34E-8	4.94E-8	2.29E-7	1.89E-6	5.35E-6
P15.	$NO_3(g) \rightarrow NO_3(p)$	8.89E-6	1.88E-5	1.03E-4	8.67E-4	2.47E-3
P16.	$PAN(g) \rightarrow PAN(p)$	7.76E-8	1.64E-7	7.60E-7	6.28E-6	1.78E-5
P17.	$O_3(g) \rightarrow O_3(p)$	1.23E-7	2.60E-7	1.21E-6	9.97E-6	2.82E-5
P18.	$OH(g) \rightarrow OH(p)$	8.39E-6	1.78E-5	9.65E-5	8.11E-4	2.31E-3
P19.	$CH_3O_2(g) \rightarrow CH_3O_2(p)$	5.49E-6	1.16E-5	5.98E-5	5.00E-4	1.42E-3
P20.	$CH_3OH(g) \rightarrow CH_3OH(p)$	4.61E-6	9.74E-6	4.95E-5	4.13E-4	1.17E-3
P21.	$DMS(g) \rightarrow DMS(p)$	6.86E-6	1.45E-5	7.68E-5	6.43E-4	1.83E-3
P22.	$DMSO(g) \rightarrow DMSO(p)$	6.21E-6	1.31E-5	6.86E-5	5.74E-4	1.63E-3
P23.	$DMSO2(g) \rightarrow DMSO2(p)$	6.06E-6	1.28E-5	6.68E-5	5.58E-4	1.59E-3
P24.	$CH_3OOH(g) \rightarrow CH_3OOH(p)$	3.55E-7	7.49E-7	3.50E-6	2.89E-5	8.18E-5
P25.	$CH_3C(O)OOH(g) \rightarrow CH_3C(O)OOH(p)$	2.83E-7	5.96E-7	2.78E-6	2.30E-5	6.50E-5
P26.	$N_2O_5(g) \rightarrow 2HNO_3(p)$	3.87E-5	8.34E-5	7.07E-4	6.23E-3	1.83E-2
P27.	$NTR(g) \rightarrow NTR(p)$	1.62E-10	3.41E-10	1.58E-9	1.31E-8	3.69E-8
P28.	$HCl(g) \rightarrow HCl(p)$	1.17E-5	2.48E-5	1.35E-4	1.13E-3	3.22E-3
P29.	$Cl(g) \rightarrow Cl(p)$	4.30E-8	9.08E-8	4.21E-7	3.48E-6	9.84E-6

Read 2.46E-6 as 2.46 x 10-6.

Table 4. Initial atmospheric conditions in the model simulations.

## (a) Gas and aqueous-phase concentrations (in ppb and M, respectively)

Species	Remote	Marine	Rural	Urban	Heavily-polluted
O3	20.0	30.0	40.0	60.0	80.0
H2O2	1.0	1.0	1.0	1.0	1.0
NO	7.5 x 10 <sup>-2</sup>	7.5 x 10 <sup>-1</sup>	1.5	10.0	100.0
NO2	2.5 x 10 <sup>-2</sup>	2.5 x 10 <sup>-1</sup>	0.5	1.0	10.0
SO2	0.6	0.6	2.0	5.0	10.0
DMS	4.0 x 10 <sup>-2</sup>	4.0 x 10 <sup>-2</sup>	0.0	0.0	0.0
НСНО	0.2	1.8	0.2	8.3	27.5
ALD2	0.0	0.0	8.3 x 10 <sup>-2</sup>	11.7	39.0
C2H6	0.0	1.9	1.58	9.08	30.3
PAR	0.0	3.1	9.8	177.9	593.1
OLE	0.0	5.6 x 10 <sup>-1</sup>	0.3	13.2	44.0
ETH	0.0	1.9 x 10 <sup>-1</sup>	0.6	11.6	38.5
TOL	0.0	3.6 x 10 <sup>-2</sup>	5.3 x 10 <sup>-1</sup>	6.0	19.9
XYL	0.0	0.0	7.5 x 10 <sup>-2</sup>	3.2	10.5
ISOP	0.0	0.0	2.4 x 10 <sup>-1</sup>	0.5	2.0
CO	80.0	100.0	120.0	150.0	300.0
CH4	$1.7 \times 10^3$	$1.7 \times 10^3$	$1.6 \times 10^3$	$1.7 \times 10^3$	$1.7 \times 10^3$
CO2	$3.4 \times 10^5$				
NH3	0.1	0.5	0.5	0.5	0.5
HCl	0.5	0.5	0.1	0.0	0.0
Fe(III)(1)	4.0 x 10 <sup>-7</sup>				
Mn(II)(l)	2.0 x 10 <sup>-7</sup>				
NaCl(1)	1.0 x 10 <sup>-6</sup>	5.0 x 10 <sup>-7</sup>	0.0	0.0	0.0

## (b) Cloud, aerosol and physical parameters

Model parameters	Remote	Marine	Rural	Urban	Heavily-polluted
Cloud					
droplet radius (µm)	10	10	10	10	10
volumetric water content	4.0 x 10 <sup>-7</sup>				
Aerosol <sup>§</sup>					
number (cm <sup>-3</sup> )	$1.26 \times 10^2$	$3.4 \times 10^2$	6.65 x 10 <sup>3</sup>	1.06 x 10 <sup>5</sup>	$2.12 \times 10^6$
mode 1 radius (µm)	5.00 x 10 <sup>-3</sup>	5.00 x 10 <sup>-3</sup>	7.39 x 10 <sup>-3</sup>	7.00 x 10 <sup>-3</sup>	6.50 x 10 <sup>-3</sup>
Log σ	0.204	0.204	0.225	0.255	0.241
number (cm <sup>-3</sup> )	2.25 x 10 <sup>1</sup>	6.00 x 10 <sup>1</sup>	1.47 x 10 <sup>2</sup>	3.20 x 10 <sup>4</sup>	3.70 x 10 <sup>4</sup>
mode 2 radius (µm)	3.55 x 10 <sup>-2</sup>	3.55 x 10 <sup>-2</sup>	2.69 x 10 <sup>-2</sup>	2.70 x 10 <sup>-2</sup>	3.1 x 10 <sup>-2</sup>
Log σ	0.301	0.301	0.557	0.334	0.297
number (cm <sup>-3</sup> )	1.50 x 10 <sup>0</sup>	$3.10 \times 10^{0}$	1.99 x 10 <sup>3</sup>	5.4 x 10 <sup>0</sup>	4.9 x 10 <sup>0</sup>
mode 3 radius (µm)	3.10 x 10 <sup>-1</sup>	3.10 x 10 <sup>-1</sup>	4.19 x 10 <sup>-2</sup>	4.3 x 10 <sup>-1</sup>	5.4 x 10 <sup>-1</sup>
Log σ	0.431	0.431	0.266	0.344	0.328
number (cm <sup>-3</sup> )	$1.50 \times 10^2$	$4.03 \times 10^2$	8.78 x 10 <sup>3</sup>	1.38 x 10 <sup>5</sup>	2.15 x 10 <sup>6</sup>
total surface (µm <sup>2</sup> cm <sup>-3</sup> )	1.39 x 10 <sup>1</sup>	2.93 x 10 <sup>1</sup>	$1.36 \times 10^2$	$1.13 \times 10^3$	$3.27 \times 10^3$
volume(µm <sup>3</sup> cm <sup>-3</sup> )	1.56 x 10 <sup>1</sup>	3.23 x 10 <sup>1</sup>	2.28 x 10 <sup>1</sup>	6.85 x 10 <sup>1</sup>	8.95 x 10 <sup>1</sup>
Temperature (K)	280	280	280	280	280
Relative humidity%	80	80	80	80	80

<sup>§</sup> Aerosol distributions for heavily polluted, urban and marine conditions are based on those of the urban and freeway, urban average, and marine surface background conditions described in Whitby, 1978. Aerosol distribution for rural condition is from Jaenicke, 1993. Aerosol distribution for remote marine condition is assumed to have the same geometric number mean diameter and geometric standard deviation as those of the marine condition, with the number concentration data for remote oceans from Fitzerald, 1991.

Table 5. 1-hr average concentrations (ppb) of gas-phase species, total sulfate (TSO4=) and total nitrate (TNO3-) in all phases under various conditions. The values in parentheses indicate the percent changes in presence of clouds and/or aerosols relative to gas only conditions.

Condition	NO	NO <sub>2</sub>	HNO <sub>3</sub>	PAN	SO <sub>2</sub>	H <sub>2</sub> SO <sub>4</sub>	НСНО	ISOP	ОН	но2	H <sub>2</sub> O <sub>2</sub>	03	TSO4=	TNO3
Remote														
Gas	3.79E-2	4.26E-2	1.66E-2		5.82E-1	2.19E-2	2.63E-1		1.484E-4	1.09E-2	1.03E+0	2.05E+1	2.19E-2	1.66E-2
Cloud	4.98E-2	4.16E-2	8.93E-6		1.24E-2	2.88E-6	1.13E-1		8.83E-5	2.22E-3	1.04E-1	2.00E+1	5.90E-1	7.72E-3
	(31.5)	(-2.3)	(-99.9)		(-97.9)	(-100.0)	(-57.0)		(-54.5)	(-79.6)	(-89.9)	(-2.4)	(2671.9)	(-53.6)
Aerosol	3.80E-2	4.26E-2	1.64E-2		5.82E-1	2.18E-2	2.61E-1		1.93E-46	1.08E-3	9.81E-1	2.05E+1	2.30E-2	1.65E-2
	(0.3)	(-0.1)	(-1.1)		(-0.1)	(-0.5)	(-0.6)		(-0.7)	(-1.1)	(-4.8)	(-0.1)	(4.9)	(-0.4)
Cloud+	4.98E-2	4.16E-2	8.91E-6		1.25E-2	2.87E-6	1.12E-1		8.83E-5	2.22E-3	1.03E-1	2.00E+1	5.90E-1	7.74E-3
Aerosol	(31.4)	(-2.3)	(-99.9)		(-97.9)	(-100.0)	(-57.2)		(-54.6)	(-79.7)	(-90.0)	(-2.4)	(2672.0)	(-53.6)
Marine														
Gas	1.99E-1	3.14E-1	2.87E-1	1.58E-1	5.67E-1	4.08E-2	1.39E+0		3.54E-4	1.11E-2	9.73E-1	3.37E+1	4.08E-2	2.87E-1
Cloud	3.53E-1	4.37E-1	1.73E-4	4.40E-2	5.87E-3	3.28E-6	1.04E+0		1.63E-4	2.73E-3	1.42E-1	3.10E+1	5.96E-1	1.47E-1
	(80.3)	(40.7)	(-99.9)	(-72.7)	(-99.0)	(-100.0)	(-25.3)		(-53.7)	(-75.2)	(-85.4)	(-7.7)	(1401.1)	(-49.1)
Aerosol	2.01E-1	3.15E-1	2.81E-1	1.56E-1	5.66E-1	4.04E-2	1.37E+0		3.50E-4	1.08E-2	8.77E-1	3.36E+1	4.29E-2	2.86E-1
	(1.2)	(0.4)	(-2.1)	(-1.2)	(-0.3)	(-1.1)	(-1.3)		(-1.0)	(-2.4)	(-9.8)	(-0.2)	(5.1)	(-0.5)
Cloud+	3.53E-1	4.37E-1	1.73E-4	4.38E-2	5.99E-3	3.19E-6	1.02		1.63E-4	2.71E-3	1.40E-1	3.10E+1	5.96E-1	1.47E-1
Aerosol	(80.4)	(40.6)	(-99.9)	(-72.8)	(-99.0)	(-100.0)	(-26.2)		(-53.8)	(-75.4)	(-85.7)	(-7.9)	(1401.0)	(-49.1)
Rural	` /			` '			` /			` /	. ,			` /
Gas	4.53E-1	7.94E-1	4.23E-1	2.04E-1	1.91E-1	8.57E-2	8.55E-1	1.15E-2	2.91E-4	5.50E-3	9.2E-1	4.27E+1	8.57E-2	4.23E-1
Cloud	5.56E-1	8.98E-1	4.07E-4	1.40E-1	4.76E-1	5.29E-5	4.43E-1	2.81E-2	1.83E-4	2.63E-3	1.19E-2	4.13E+1	1.52E+0	3.18E-1
	(23.5)	(13.5)	(-99.9)	(-31.0)	(-75.1)	(-99.9)	(-48.2)	(190.4)	(-37.0)	(-51.7)	(-98.7)	(-3.2)	(1751.6)	(-24.9)
Aerosol	4.73E-1	8.10E-1	3.75E-1	1.92E-1	1.89E+0	7.98E-2	7.98E-1	1.30E-2	2.66E-4	4.73E-3	5.16E-1	4.22E+1	1.19E-1	4.05E-1
	(4.5)	(2.1)	(-11.0)	(-5.7)	(-1.1)	(-6.8)	(-6.6)	(16.5)	(-8.6)	(-13.5)	(-43.9)	(-1.1)	(39.1)	(-4.2)
Cloud+	5.61E-1	8.99E-1	3.98E-4	1.38E-1	4.84E-1	5.20E-5	4.23E-1	2.91E-2	1.79E-4	2.53E-3	1.10E-2	4.10E+1	1.52E+0	3.16E-1
Aerosol	(24.7)	(13.6)	(-99.9)	(-32.3)	(-74.7)	(-99.9)	(-50.5)	(202.7)	(-38.3)	(-53.4)	(-98.8)	(-3.9)	(1747.4)	(-25.4)
Urban	(2)	(15.0)	( , , , , ,	(52.5)	( , ,	( , , , , )	(20.2)	(20217)	(20.2)	( 55)	( ) 0.0)	(3.7)	(17.77.1)	(2011)
Gas	1.82E+0	5.19E+0	9.91E-1	2.44E+0	4.93E+0	7.24E-2	1.04E+1	1.57E-1	9.15E-5	6.06E-3	9.56E-1	7.18E+1	7.24E-2	9.91E-1
Cloud	2.07E+0	5.52E+0	1.15E-3	1.98E+0	2.80E+0	1.52E-4	7.97E+0	1.88E-1	7.35E-5	4.17E-3	3.06E-3	6.82E+1	2.16E+0	9.84E-1
Cloud	(14.7)	(6.6)	(-99.9)	(-18.5)	(-43.2)	(-99.8)	(-23.6)	(20.8)	(-19.7)	(-30.4)	(-99.7)	(-5.0)	(2977.7)	(-0.7)
Aerosol	2.14E+0	5.44E+0	4.76E-1	1.91E+0	4.41E+0	5.35E-2	6.07E+0	1.90E-1	6.70E-5	3.58E-3	1.23E-2	6.50E+1	8.28E-1	9.55E-1
71010301	(19.2)	(5.1)	(-50.9)	(-21.0)	(-10.4)	(-25.4)	(-41.7)	(22.6)	(-26.7)	(-40.1)	(-98.7)	(-9.3)	(1051.6)	(-3.3)
Cloud+	2.29E+0	5.60E+0	9.53E-4	1.69E+0	2.61E+0	1.16E-4	5.13E+0	2.09E-1	6.06E-5	3.00E-3	1.94E-3	6.31E+1	2.51E-1	1.01E+0
Aerosol	(27.6)	(8.4)	(-99.9)	(-30.4)	(-47.0)	(-99.8)	(-50.7)	(35.7)	(-33.9)	(-49.7)	(-99.9)	(-12.0)	(3472.2)	(2.1)
Heavily-	(27.0)	(0.4)	( )).))	( 30.4)	( 47.0)	( )).0)	( 30.7)	(33.1)	( 33.7)	( 47.1)	( )).))	(12.0)	(3472.2)	(2.1)
Polluted														
Gas	2.41E+1	7.51E+1	5.52E+0	4.02E+0	9.93E+1	6.99E-2	2.85E+1	1.04E+0	4.30E-5	8.63E-4	9.45E-1	8.35E+1	6.99E-2	5.52E+0
Cloud	2.47E+1 2.47E+1	7.31E+1 7.32E+1	7.46E-3	3.71E+0	9.93E+1 8.17E+1	3.87E-4	2.83E+1 2.39E+1	1.04E+0 1.08E+0	3.99E-5	7.75E-4	3.25E-5	7.97E+1	0.99E-2 1.73E+0	7.37E+0
Ciouu	(2.3)	(-2.5)	(-99.9)	3.71E+0 (-7.6)	(-17.7)	3.87E-4 (-99.4)	(-16.2)	(4.0)	3.99E-3 (-6.8)	(-10.4)	(-100.0)	(-4.6)	(2463.3)	(33.3)
Agrosol	2.78E+1	(-2.5) 6.87E+1	(-99.9) 1.01E+0	(-7.6) 2.65E+0	7.24E+1	(-99.4) 3.77E-2	6.73E+0	1.22E+0	(-0.8) 2.53E-5	(-10.4) 4.09E-4	(-100.0) 2.0E-5	6.63E+1	(2403.3) 4.00E+0	8.26E+0
Aerosol		6.87E+1 (-8.5)		2.03E+0 (-33.2)	(-27.1)	3.77E-2 (-45.0)		(18.1)		4.09E-4 (-52.4)	2.0E-5 (-100.0)		4.00E+0 (5670.9)	8.26E+0 (51.0)
Cloud	(15.6)		(-80.7)				(-76.4)		(-40.9)			(-20.4)		
Cloud+	2.78E+1 (15.6)	6.84E+1	4.55E-3 (-99.9)	2.68E+0	5.99E+1 (-39.7)	2.05E-4 (-99.7)	7.18E+0	1.22E+0	2.63E-5	4.22E-4	1.59E-5 (-100.0)	6.60E+1 (-20.8)	4.96E+0	9.85E+0
Aerosol	(13.0)	(-8.9)	(-99.9)	(-32.7)	(-39./)	(-99./)	(-74.8)	(17.8)	(-38.7)	(-50.8)	(-100.0)	(-20.8)	(7135.1)	(79.8)

Table 6. Predicted photochemical indicators under various polluted atmospheric conditions and the changes in the presence of clouds and/or aerosols relative to the base value under clear air conditions at the end of the 2-hr simulation.

Indicators	Scenarios			Cor	nditions		
		]	Rural		Jrban	Heavil	y-polluted
		Indicator value	Relative change	Indicator value	Relative change	Indicator value	Relative change
$NO_y$	Gas	2.00E+0		1.10E+1		1.10E+2	
,	Cloud	1.59E+0	-2.05E-1	9.71E+0	-1.17E-1	9.99E+1	-9.18E-2
	Aerosol	1.95E+0	-2.50E-2	1.01E+1	-8.18E-2	9.78E+1	-1.11E-1
	Cloud+Aerosol	1.59E+0	-2.05E-1	9.63E+0	-1.25E-1	9.64E+1	-1.24E-1
O <sub>3</sub> /NO <sub>V</sub>	Gas	2.19E+1		7.05E+0		8.31E-1	
,	Cloud	2.64E+1	2.06E-1	7.49E+0	6.24E-2	8.64E-1	3.97E-2
	Aerosol	2.21E+1	9.10E-3	6.72E+0	-4.68E-2	6.86E-1	-1.75E-1
	Cloud+Aerosol	2.61E+1	1.92E-1	6.81E+0	-3.40E-2	6.94E-1	-1.65E-1
O <sub>3</sub> /NO <sub>z</sub>	Gas	4.62E+1		1.47E+1		6.48E+0	
	Cloud	1.52E+2	2.29E+0	2.26E+1	5.37E-1	1.41E+1	1.18E+0
	Aerosol	5.08E+1	9.96E-2	1.91E+1	2.99E-1	1.32E+1	1.04E+0
	Cloud+Aerosol	1.53E+2	2.31E+0	2.45E+1	6.67E-1	1.60E+1	1.47E+0
O <sub>3</sub> /HNO <sub>3</sub>	Gas	8.01E+1		5.97E+1		1.22E+1	
	Cloud	1.18E+5	1.47E+3	6.99E+4	1.17E+3	1.15E+4	9.42E+2
	Aerosol	9.11E+1	1.37E-1	1.29E+2	1.16E+0	6.92E+1	4.67E+0
	Cloud+Aerosol	1.20E+5	1.50E+3	7.67E+4	1.28E+3	1.59E+4	9.50E+3
H <sub>2</sub> O <sub>2</sub> /NO <sub>y</sub>	Gas	4.50E-1		8.64E-2		8.45E-3	
•	Cloud	8.06E-3	-9.82E-1	4.04E-4	-9.95E-1	3.23E-7	-1.00E+0
	Aerosol	2.15E-1	-5.22E-1	4.16E-4	-9.95E-1	1.23E-7	-1.00E+0
	Cloud+Aerosol	7.29E-3	-9.84E-1	2.29E-4	-9.97E-1	1.46E-7	-1.00E+0
H <sub>2</sub> O <sub>2</sub> /HNO <sub>3</sub>	Gas	1.64E+0		1.00E+0		1.24E-1	
	Cloud	3.62E+1	2.11E+1	3.77E+0	2.77E+0	4.31E-3	-9.65E-1
	Aerosol	8.84E-1	-4.61E-1	8.00E-3	-9.92E-1	1.24E-5	-9.99E-1
	Cloud+Aerosol	3.34E+1	1.94E+1	2.58E+0	1.58E+0	3.35E-3	-9.73E-1
HCHO/NO <sub>y</sub>	Gas	4.64E-1		9.49E-1		2.62E-1	
•	Cloud	3.03E-1	-3.47E-1	8.49E-1	-1.05E-1	2.44E-1	-6.87E-2
	Aerosol	4.38E-1	-5.60E-2	5.47E-1	-4.24E-1	5.24E-2	-8.00E-1
	Cloud+Aerosol	2.85E-1	-3.86E-1	4.96E-1	-4.77E-1	5.84E-2	-7.77E-1
(HNO <sub>3</sub> /H <sub>2</sub> O <sub>2</sub> )/	Gas	2.74E-1		1.18E-1		6.80E-2	
(NO <sub>X</sub> /ROG)	Cloud	2.23E-4	-9.99E-1	1.04E-4	-9.99E-1	7.50E-5	-9.99E-1
-	Aerosol	2.43E-1	-1.13E-1	5.20E-2	-5.59E-1	9.92E-3	-8.54E-1
	Cloud+Aerosol	2.18E-4	-9.99E-1	8.87E-5	-9.99E-1	4.37E-5	-9.99E-1

Table 7. 1-hr average sensitivities of  $O_3$  with respect to gas-phase reaction rate constants under clear air conditions.

	Re	mote	M	arine	]	Rural	U	Irban	Heavil	y-polluted
Rank	Rxn. #	$\overline{S}$ kl	Rxn. #	$\overline{S}_{kl}$	Rxn. #	$\frac{\overline{S}}{S}$ kl	Rxn. #	$\frac{\overline{S}}{S}$ kl	Rxn. #	$\overline{S}$ kl
1	G28	2.01E-2	G1	6.03E-2	G11	4.67E-2	G38	1.20E-1	G3	-2.58E-1
2	G1	1.86E-2	G3	-4.92E-2	G10	-4.67E-2	G43	-1.14E-1	G1	2.77E-1
3	G3	-1.29E-2	G28	4.40E-2	<b>G</b> 9	4.47E-2	G1	1.10E-1	G38	1.85E-1
4	G116	6.44E-3	G26	-3.25E-2	G1	4.13E-2	G3	-1.06E-1	G26	-1.75E-1
5	G117	4.38E-3	G11	2.26E-2	G26	-4.10E-2	G57	8.05E-2	G57	1.23E-1
6	G118	-4.05E-3	G10	-2.26E-2	G3	-3.87E-2	G26	-6.34E-2	G43	-8.70E-2
7	G26	-3.94E-3	G9	2.17E-2	G36	1.54E-2	G11	5.12E-2	G45	5.07E-2
8	G33	-3.33E-3	G38	2.10E-2	G116	1.50E-2	G10	-5.12E-2	G47	-4.11E-2
9	G32	-2.81E-3	G116	1.97E-2	G28	1.32E-2	G9	4.90E-2	G46	4.11E-2
10	G13	-2.62E-3	G36	1.46E-2	G38	1.04E-2	G47	-4.60E-2	G58	4.01E-2
11	G34	2.29E-3	G43	-9.41E-3	G61	8.65E-3	G46	4.60E-2	G61	3.93E-2
12	G12	-1.50E-3	G33	-8.24E-3	G34	8.48E-3	G61	3.31E-2	G69	3.79E-2
13	G36	1.33E-3	G32	-6.95E-3	G47	-7.96E-3	G45	2.99E-2	G52	3.38E-2
14	G38	1.12E-3	G47	-6.72E-3	G46	7.94E-3	G69	2.82E-2	G39	-2.74E-2
15	G29	-7.65E-4	G46	6.61E-3	G43	-6.15E-3	G28	2.35E-2	G11	2.61E-2
16	G30	7.48E-4	G117	6.29E-3	G52	6.14E-3	G52	2.23E-2	G10	-2.61E-2
17	G97	-5.59E-4	G34	6.03E-3	G74	5.21E-3	G58	2.08E-2	G9	2.49E-2
18	G89	-5.58E-4	G118	-5.64E-3	G57	3.72E-3	G74	1.74E-2	G74	1.71E-2
19	G11	5.36E-4	G37	4.84E-3	G76	3.52E-3	G39	-1.68E-2	G28	1.50E-2
20	G10	-5.36E-4	G39	-3.72E-3	G29	-3.20E-3	G29	-1.47E-2	G29	-1.46E-2
21	G85	3.70E-4	G29	-3.58E-3	G30	3.12E-3	G30	1.42E-2	G30	1.44E-2
22	G9	3.07E-4	G30	3.43E-3	G63	2.26E-3	G37	6.24E-3	G4	-1.42E-2
23	G122	-2.55E-4	G13	-3.04E-3	G37	2.05E-3	G34	6.14E-3	G63	1.11E-2
24	G22	-2.25E-4	G57	2.88E-3	G22	-1.99E-3	G63	5.54E-3	G76	9.99E-3
25	G23	1.88E-4	G61	2.55E-3	G69	1.90E-3	G36	4.17E-3	G2	8.87E-3
26	G39	-9.66E-5	G52	2.35E-3	G33	-1.85E-3	G76	4.16E-3	G37	8.84E-3
27	G82	9.60E-5	G22	-1.65E-3	G73	-1.84E-3	G116	3.14E-3	G65	-4.78E-3
28	G37	-8.24E-5	G12	-1.55E-3	G23	1.78E-3	G73	-2.89E-3	G64	4.78E-3
29	G86	-6.43E-5	G23	1.44E-3	G32	-1.56E-3	G64	2.31E-3	G7	-4.60E-3
30	G88	6.28E-5	G35	7.64E-4	G13	-1.51E-3	G65	-2.30E-3	G71	4.47E-3

Table 8. 1-hr average sensitivities of odd hydrogen  $R_xO_y$  with respect to gas-phase reaction rate constants under clear air conditions.

	Re	emote	M	Iarine		Rural	J	Jrban	Heavi	ly-polluted
Rank	Rxn.#	$\overline{S}_{kl}$	Rxn. #	$\overline{\overline{S}}$ kl	Rxn. #	$\overline{S}_{kl}$	Rxn.#	$\overline{S}_{kl}$	Rxn. #	$\overline{S}_{kl}$
1	G9	3.65E-1	G9	5.16E-1	G11	1.03E+0	G38	8.42E-1	G38	7.17E-1
2	G11	3.52E-1	G11	5.05E-1	G10	-1.03E+0	G28	-5.80E-1	G28	-6.53E-1
3	G10	-3.52E-1	G10	-5.05E-1	G9	1.03E+0	G43	-5.69E-1	G26	-6.21E-1
4	G118	-1.80E-1	G38	2.90E-1	G28	-4.99E-1	G11	4.03E-1	G57	4.48E-1
5	G33	-1.62E-1	G116	1.72E-1	G26	-4.27E-1	G10	-4.03E-1	G43	-3.21E-1
6	G116	1.49E-1	G33	-1.61E-1	G116	2.54E-1	G9	4.02E-1	G117	-2.54E-1
7	G32	-1.37E-1	G32	-1.36E-1	G38	2.46E-1	G57	3.60E-1	G58	2.34E-1
8	G34	1.28E-1	G117	-1.09E-1	G36	2.02E-1	G1	-3.20E-1	G3	2.34E-1
9	G117	-1.22E-1	G34	1.06E-1	G34	1.63E-1	G3	2.88E-1	G1	-2.28E-1
10	G28	8.71E-2	G118	-9.78E-2	G1	-1.61E-1	G26	-2.77E-1	G45	2.09E-1
11	G36	-6.94E-2	G37	-8.74E-2	G3	1.41E-1	G117	-2.48E-1	G69	1.72E-1
12	G38	5.26E-2	G39	-6.87E-2	G117	-1.38E-1	G47	-2.45E-1	G39	-1.65E-1
13	G37	-1.95E-2	G26	-6.81E-2	G74	1.12E-1	G69	2.26E-1	G47	-1.57E-1
14	G1	-1.50E-2	G47	-5.06E-2	G47	-1.06E-1	G46	2.24E-1	G46	1.42E-1
15	G35	-1.26E-2	G36	4.16E-2	G46	9.31E-2	G45	2.10E-1	G9	1.23E-1
16	G26	-1.25E-2	G28	-4.06E-2	G61	9.13E-2	G58	1.79E-1	G61	1.21E-1
17	G3	1.03E-2	G46	3.73E-2	G43	-8.30E-2	G74	1.72E-1	G11	1.21E-1

Table 9. 1-hr average sensitivities of gas-phase species and photochemical indicators with respect to temperature (Temp.) and relative humidity (R.H.) under various clear air conditions.

Species or	Model parameters			Atmospheric condition	S	
indicators	•	Remote	Marine Ru	ıral Urban	Heavily-polluted	
NO2	Temp.	2.37E+0	-1.81E+0	-4.94E+0	-2.65E+0	9.79E-1
	R.H.	-3.96E-2	-2.10E-1	-3.22E-1	-1.34E-1	-7.40E-2
SO2	Temp.	-1.47E-1	-3.33E-1	-7.44E-1	-1.50E-1	-4.97E-2
	R.H.	-9.02E-3	-1.54E-2	-3.13E-2	-3.73E-3	-8.01E-3
HNO3	Temp.	6.62E+0	6.35E+0	1.42E+1	9.66E+0	6.30E+0
	R.H.	3.08E-1	2.02E-1	5.80E-1	2.19E-1	1.36E-1
H2SO4	Temp.	5.18E+0	6.21E+0	1.67E+1	1.03E+1	7.06E+0
	R.H.	2.99E-1	2.62E-1	7.01E-1	2.54E-1	1.13E-1
OH	Temp.	5.48E+0	5.79E+0	1.60E+1	1.02E+1	7.47E+0
	R.H.	2.87E-1	2.18E-1	6.69E-1	2.62E-1	1.27E-1
HO2	Temp.	7.79E+0	1.37E+1	2.86E+1	1.92E+1	1.15E+1
	R.H.	1.20E-1	3.37E-1	1.00E+0	3.95E-1	1.20E-1
PAN	Temp.		5.71E+0	1.33E+1	8.23E+0	4.51E+0
	R.H.		3.75E-1	6.91E-1	2.76E-1	1.18E-1
O3	Temp.	3.48E-2	7.29E-1	1.26E+0	2.29E+0	1.12E+0
	R.H.	-2.80E-3	1.43E-2	4.49E-2	4.96E-2	2.52E-2
NOy	Temp.			1.41E-1	3.02E-1	9.93E-2
	R.H.			-8.60E-3	-3.45E-3	-3.66E-3
O3/NOz	Temp.			-1.28E+1	-6.93E+0	-4.75E+0
	R. H.			-5.67E-1	-2.10E-1	-1.02E-1
H2O2/HNO3	Temp.			-1.43E+1	-9.22E+0	-6.36E+0
	R.H.			-5.76E-1	-1.99E-1	-1.37E-1

Table 10. 1-hr average sensitivities of  $O_3$  with respect to aqueous-phase reaction rate constants under various cloudy conditions.

	Re	mote	M	arine	R	tural	U	Irban	Heavil	y-polluted
Rank	Rxn.#	$\overline{S}$ kl	Rxn. #	$\overline{S}_{kl}$	Rxn. #	$\overline{S}$ $_{kl}$	Rxn. #	$\overline{S}$ kl	Rxn. #	$\overline{S}$ kl
1	L13	-3.14E-3	L7	-7.76E-3	L7	-7.08E-3	L7	-1.22E-2	L109	6.13E-3
2	L7	-5.96E-4	L13	-4.96E-3	L13	-2.46E-3	L13	-2.52E-3	L75	-1.31E-3
3	L74	-3.86E-5	L75	6.78E-4	L74	-7.21E-4	L105	-1.93E-3	L105	-6.65E-4
4	L50	-3.70E-5	L74	-2.19E-4	L75	6.45E-4	L6	-8.36E-4	L112	-6.62E-4
5	L75	3.29E-5	L6	-1.82E-4	L82	-4.52E-4	L112	-7.60E-4	L7	-4.71E-4
6	L16	-2.13E-5	L105	-1.02E-4	L6	-3.29E-4	L109	7.35E-4	L13	-3.24E-4
7	L6	-1.92E-5	L50	-8.52E-5	L105	-3.03E-4	L82	-4.93E-4	L74	-2.60E-4
8	L105	1.65E-5	L1	5.20E-5	L79	-1.92E-4	L50	4.84E-4	L50	1.30E-4
9	L64	-1.17E-5	L112	-4.64E-5	L85	1.55E-4	L74	-4.38E-4	L44	1.16E-4
10	L49	-9.16E-6	L57	3.83E-5	L57	9.48E-5	L114	-3.36E-4	L32	1.16E-4
11	L5	-8.91E-6	L19	-3.37E-5	L80	9.01E-5	L85	1.90E-4	L99	8.78E-5
12	L1	6.89E-6	L18	-3.31E-5	L3	-8.88E-5	L79	-1.73E-4	L114	-8.24E-5
13	L73	-6.85E-6	L20	3.27E-5	L50	8.48E-5	L99	1.60E-4	L82	-8.10E-5
14	L3	-5.21E-6	L49	-2.49E-5	L81	8.19E-5	L18	-1.55E-4	L73	-6.78E-5
15	L57	-4.38E-6	L114	-2.28E-5	L77	6.93E-5	L46	-9.43E-5	L6	-5.58E-5
16	L82	2.16E-6	L16	-1.88E-5	L91	6.36E-5	L91	-8.47E-5	L91	-4.63E-5
17	L77	1.81E-6	L64	-1.61E-5	L18	-5.40E-5	L93	-7.91E-5	L85	4.13E-5
18	L4	-1.61E-6	L77	1.53E-5	L19	-4.74E-5	L19	-7.11E-5	L93	-4.02E-5
19	L23	-1.52E-6	L82	-1.36E-5	L109	4.43E-5	L59	7.08E-5	L31	3.95E-5
20	L21	-1.51E-6	L25	-1.25E-5	L20	4.40E-5	L77	7.04E-5	L46	-3.79E-5
21	L18	-1.38E-6	L5	1.25E-5	L96	-3.58E-5	L106	-6.82E-5	L59	2.79E-5
22	L22	1.35E-6	L73	-1.17E-5	L73	-3.26E-5	L80	6.29E-5	L79	-2.38E-5
23	L19	-1.34E-6	L3	-1.14E-5	L106	3.19E-5	L20	6.09E-5	L18	-1.78E-5
24	L20	1.32E-6	154	1.11E-5	L25	-3.10E-5	L73	-5.66E-5	L43	-1.45E-5
25	L85	-1.19E-6	L79	-1.07E-5	L24	2.86E-5	L57	2.89E-5	L97	-1.31E-5

Table 11. 1-hr average sensitivities of aqueous-phase S(IV) and S(VI) with respect to aqueous-phase reaction rate constants under various cloudy conditions.

(a) S(IV)

	Re	mote	M	Iarine		Rural	J	Jrban	Heavi	ly-polluted
Rank	Rxn.#	$\overline{S}_{kl}$	Rxn. #	$\overline{S}$ kl	Rxn. #	$\overline{S}_{kl}$	Rxn.#	$\overline{S}$ kl	Rxn. #	$\overline{S}_{kl}$
1	L75	-3.52E+0	L75	-4.52E+0	L75	2.95E-1	L105	-1.66E-1	L105	-2.84E-2
2	L105	-2.69E-1	L105	-4.23E-1	L105	-2.14E-1	L75	6.17E-2	L112	-1.58E-2
3	L7	1.32E-1	L7	1.52E-1	L82	6.17E-2	L77	-2.27E-2	L109	-1.44E-2
4	L5	5.65E-2	L77	-9.33E-2	L77	-5.53E-2	L50	2.26E-2	L77	-1.23E-2
5	L77	-5.50E-2	L112	-6.09E-2	L91	-5.27E-2	L82	2.07E-2	L50	1.02E-2
6	L79	-4.14E-2	L74	-4.08E-2	L80	-5.23E-2	L112	-1.91E-2	L75	6.56E-3
7	L103	-2.15E-2	L79	-3.51E-2	L81	-3.95E-2	L79	-1.82E-2	L91	-6.32E-3
8	L50	-1.88E-2	L106	-3.03E-2	L79	-3.89E-2	L80	-1.43E-2	L80	-5.75E-3
9	L13	1.70E-2	L114	-2.10E-2	L74	-3.51E-2	L7	1.38E-2	L82	4.91E-3
10	L106	-8.54E-3	L50	1.99E-2	L7	3.26E-2	L91	-1.08E-2	L108	-3.39E-3

(b) S(VI)

	Re	mote	M	arine		Rural	U	rban	Heavil	y-polluted
Rank	Rxn.#	$\overline{S}$ kl	Rxn. #	$\overline{S}_{kl}$	Rxn. #	$\frac{\overline{S}}{S}$ kl	Rxn. #	$\overline{S}_{kl}$	Rxn. #	$\overline{S}$ kl
1	L75	8.60E-2	L75	7.22E-2	L75	-1.14E-1	L105	2.47E-1	L105	1.40E-1
2	L105	6.41E-3	L105	6.56E-3	L105	8.49E-2	L75	-7.79E-2	L109	7.18E-2
3	L7	-3.45E-3	L7	-3.93E-3	L82	-2.47E-2	L50	-3.86E-2	L77	6.02E-2
4	L77	1.31E-3	L112	-3.27E-3	L77	2.20E-2	L77	3.34E-2	L50	-4.44E-2
5	L5	-1.26E-3	L114	2.30E-3	L91	2.11E-2	L82	-3.13E-2	L91	3.08E-2
6	L79	9.91E-4	L50	-2.26E-3	L80	2.09E-2	L79	2.71E-2	L80	2.84E-2
7	L103	3.85E-4	L77	1.40E-3	L81	1.58E-2	L80	2.15E-2	L82	-2.46E-2
8	L50	3.22E-4	L13	1.09E-3	L79	1.55E-2	L7	-2.15E-2	L114	1.81E-2
9	L13	-2.81E-4	L74	6.06E-4	L74	1.40E-2	L91	1.61E-2	L108	1.66E-2
10	L106	2.07E-4	L106	4.92E-4	L7	-1.30E-2	L74	1.28E-2	L75	1.62E-2

Table 12. 1-hr average sensitivities of  $O_3$  with respect to equilibrium constants under various cloudy conditions. Only those sensitivities greater than 1.0E-4 are shown.

(a) Dissolution equlibrium constant (i.e., Henry's Law constant)

	Rei	mote	Ma	rine	R	ural	U1	ban	Heavily	-polluted
Rank	Rxn. #	$\overline{S}_{kl}$	Rxn. #	$\overline{S}$ kl	Rxn. #	$\overline{S}_{kl}$	Rxn. #	$\overline{S}$ $_{kl}$	Rxn. #	$\overline{S}$ $_{kl}$
1	E10	-4.38E-3	E10	-2.11E-2	E10	-1.83E-2	E10	-3.16E-2	E12	-3.01E-2
2	E17	-3.23E-3	E17	-5.28E-3	E17	-3.39E-3	E12	-1.53E-2	E13	6.61E-3
3	E11	-1.20E-4	E12	-1.51E-3	E3	-9.77E-4	E17	-3.13E-3	E3	4.25E-3
4	E18	-1.02E-4	E11	-3.36E-4	E12	-5.31E-4	E3	-2.37E-3	E10	-2.24E-3
5			E3	-2.03E-4	E18	-1.96E-4	E13	7.57E-4	E17	-6.86E-4
6			E18	-1.35E-4			E5	2.43E-4	E5	1.10E-4
7							E7	-1.43E-4		

(b) Dissociation equlibrium constant

	Remote		Marine		Rural		Urban		Heavily-polluted	
Rank	Rxn.#	$\overline{S}$ $_{kl}$	Rxn.#	$\overline{S}_{kl}$	Rxn. #	$\overline{S}$ $_{kl}$	Rxn. #	$\overline{S}_{kl}$	Rxn. #	$\overline{S}$ $_{kl}$
1	D2	-3.75E-3	D2	-1.28E-2	D2	-1.01E-2	D2	-1.55E-2	D10	4.26E-3
2			D11	-2.64E-4	D10	-9.73E-4	D10	-2.34E-3	D11	-9.25E-4
3			D10	-2.03E-4	D11	-6.63E-4	D11	-1.18E-3	D2	-9.10E-4
4							D9	7.62E-4	D9	6.65E-4
5							D4	-3.58E-4	D16	1.21E-4
6							D16	2.62E-4		

Table 13. 1-hr average sensitivities of species and photochemical indicators with respect to temperature (Temp.), relative humidity (R.H.), cloud droplet radius (Rdrop.) and cloud liquid water ( $V_c$ ) under various cloudy conditions. TSO<sub>4</sub><sup>2-</sup> and TNO<sub>3</sub><sup>-</sup> are the total sulfate and nitrate in all phases, respectively.

Species	Model parameters		A	tmospheric conditions		
		Remote	Marine Rui	al Urban	Heavily-polluted	
NO2	Temp.	2.52E+0	-6.76E-1	-2.86E+0	-2.54E+0	6.51E-1
	R.H.	-2.33E-2	-1.04E-1	-1.79E-1	-9.29E-2	-5.81E-3
	Rdrop.	4.52E-3	-7.56E-2	-4.69E-2	-2.70E-2	2.08E-3
	$V_c$	-9.55E-3	7.32E-2	6.14E-2	5.82E-2	-1.27E-3
SO2	Temp.	4.09E+1	5.08E+1	5.05E+0	3.89E+0	1.93E+(
	R.H.	-4.81E-1	-5.41E-1	-3.25E-1	-6.67E-2	-1.61E-3
	Rdrop.	1.71E-2	1.46E-1	-7.37E-2	9.00E-2	-7.94E-3
	$V_c$	-6.98E-1	-1.05E+0	-3.88E-1	-2.45E-1	-7.28E-2
HNO3	Temp.	1.81E+1	1.38E+1	1.56E+1	1.10E+1	8.28E+0
	R.H.	7.30E-1	4.06E-1	5.04E-1	1.84E-1	1.28E-1
	Rdrop.	2.23E+0	2.21E+0	2.08E+0	1.98E+0	1.91E+(
	$V_c$	-1.30E+0	-1.33E+0	-1.23E+0	-1.15E+0	-1.07E+0
H2SO4	Temp.	1.56E+1	8.11E+0	2.00E+1	1.16E+1	4.68E+0
	R.H.	9.87E-2	1.29E-1	2.96E-1	1.59E-1	5.35E-2
	Rdrop.	4.31E-1	1.05E+0	1.29E+0	1.04E+0	6.02E-1
	$V_c$	-1.16E+0	-1.29E+0	-1.43E+0	-1.11E+0	-8.45E-1
ОН	Temp.	1.82E+1	1.70E+1	2.09E+1	1.35E+1	9.32E+(
011	R.H.	7.55E-1	5.10E-1	6.84E-1	2.63E-1	1.31E-1
	Rdrop.	3.17E-1	3.76E-1	2.17E-1	8.93E-2	-5.62E-4
	$V_c$	-2.49E+0	-4.01E-1	-2.94E-1	-1.91E-1	-6.65E-2
HO2	Temp.	2.71E+1	2.41E+1	3.00E+1	2.28E+1	1.43E+1
1102	R.H.	5.06E-1	4.29E-1	7.16E-1	3.27E-1	1.23E-1
	Rdrop.	4.37E-1	4.73E-1	2.37E-1	1.36E-1	-2.92E-3
	L.W.	-6.33E-1	-6.07E-1	-4.24E-1	-3.18E-1	-1.06E-1
PAN	Temp.	0.33 <u>L</u> 1	2.34E+1	1.63E+1	1.15E+1	5.97E+(
17111	R.H.		8.89E-1	6.91E-1	2.89E-1	1.24E-1
	Rdrop.		7.46E-1	1.97E-1	9.37E-2	2.00E-3
	$V_c$		-7.41E-1	-2.58E-1	-1.95E-1	-6.30E-2
O3	Temp.	7.21E-2	6.63E-1	8.95E-1	2.56E+0	1.39E+(
03	R.H.	-2.12E-3	1.03E-2	2.22E-2	4.18E-2	2.59E-2
	Rdrop.	3.86E-3	2.16E-2	9.03E-3	1.94E-2	-1.26E-3
	$V_c$	-6.04E-3	-2.69E-2	-1.69E-2	-4.35E-2	-2.67E-2
	Temp.	-9.38E-1	-5.33E-1	-2.04E+0	-5.20E+0	-7.73E+(
TSO4 <sup>2</sup> -	*					
	R.H.	1.29E-2	1.06E-2	1.28E-1	9.82E-2	1.15E-2
	Rdrop.	3.86E-4 -9.84E-1	-3.51E-3 -9.91E-1	2.95E-2 -8.47E-1	-1.49E-1 -6.53E-1	5.95E-3
	$V_{c}$					-6.87E-1
TNO3	Temp.	1.85E+1	1.59E+1	1.88E+1	1.29E+1	9.41E+(
	R.H.	6.91E-1	4.57E-1	6.18E-1	1.99E-1	8.70E-2
	Rdrop.	2.13E-1	3.57E-1	1.59E-1	4.82E-2	-2.61E-2
	$V_C$	-1.22E+0	-1.37E+0	-1.22E+0	-1.12E+0	-1.03E+0
NOy	Temp.			-3.10E+0	-1.04E+0	-6.22E-1
•	R.H.			-1.07E-1	-2.10E-2	-6.68E-3
	Rdrop.			-2.78E-2	-5.47E-1	1.91E-3
	$V_c$			3.96E-2	1.42E-2	2.58E-3
O3/NOz	Temp.			-1.60E+1	-9.74E+0	-5.38E+0
	R.H.			-6.36E-1	-2.46E-1	-9.81E-2
	Rdrop.			-1.83E-1	-7.54E-2	-5.38E-2
	$V_c$			2.31E-1	1.51E-1	3.81E-
H2O2/HNO3	Temp.			3.86E+1	3.40E+1	1.85E+
11202/111103	R.H.			6.96E-1	4.54E-1	1.70E-
	Rdrop.			-2.09E+0	-2.41E+0	-1.91E+(
	$V_c$			7.89E-1	7.95E-1	8.15E-1

Table 14. 1-hr average sensitivity of various species concentrations with respect to uptake coefficients of individual species under the Heavily-polluted aerosol conditions. Only those sensitivities greater than 1.0E-4 are shown.

(a) Reactive nitrogen species

Rank		NO		NO2		NO3		N2O5		HNO3		PAN
1	O3	7.78E-2	NO2	-3.95E-2	O3	-1.97E-1	N2O5	-8.02E-1	HNO3	-9.01E-1	HCHO	-2.31E-1
2	HCHO	6.44E-2	O3	-2.56E-2	HCHO	-1.20E-1	O3	-2.21E-1	HCHO	-2.70E-1	PAN	-4.63E-2
3	NO2	-3.15E-2	NO	-7.21E-3	N2O5	-1.82E-2	HCHO	-1.20E-1	O3	-3.73E-2	O3	-1.73E-2
4	NO	-1.50E-2	N2O5	-3.46E-3	NO2	-1.63E-2	NO2	-5.55E-2	NO2	-1.45E-2	HO2	-1.34E-2
5	HO2	3.58E-3	HCHO	-8.01E-4	NO	1.43E-2	NO	7.03E-3	HO2	-1.44E-2	NO2	7.94E-3
6	N2O5	-1.27E-3	HNO2	-2.51E-4	HO2	-6.56E-3	HO2	-6.50E-3	HNO2	-9.57E-3	HNO2	-7.88E-3
7	HNO2	1.06E-3			HNO2	-2.39E-3	HNO2	-2.62E-3	N2O5	-7.61E-3	NO	3.63E-3
8	H2O2	3.06E-4			H2O2	-5.20E-4	H2O2	-4.96E-4	SO2	1.29E-3	H2O2	-1.43E-3
9	SO2	-2.46E-4			SO2	4.17E-4	SO2	3.97E-4	H2O2	-6.51E-4	SO2	1.05E-3
10					CH3O2	-1.69E-4	CH3O2	-1.72E-4	CH3O2	-3.16E-4	N2O5	3.34E-4
11					NO3	-1.27E-4	NH3	-1.39E-4	PAN	-2.15E-4	CH3O2	2.84E-4
12							PAN	-1.08E-4				

(b) Odd hydrogen species

Rank	(	ОН		HO2		H2O2	(	C2O3	CI	H3O2	CH3	ООН
1	НСНО	-2.86E-1	HCHO	-3.82E-1	H2O2	-2.99E+0	НСНО	-2.86E-1	HCHO	-2.72E-1	НСНО	-5.06E-1
2	O3	-2.00E-2	O3	-9.50E-2	HCHO	-6.14E-1	NO2	5.42E-2	O3	-9.07E-2	СНЗООН	-2.10E-1
3	NO2	1.91E-2	NO2	4.40E-2	O3	-1.50E-1	HO2	-1.59E-2	NO2	4.59E-2	O3	-9.38E-2
4	HO2	-1.58E-2	HO2	-2.00E-2	NO2	6.92E-2	NO	1.40E-2	NO	1.96E-2	NO2	4.55E-2
5	HNO2	-1.03E-2	NO	1.95E-2	HO2	-3.19E-2	HNO2	-9.80E-3	HO2	-1.50E-2	HO2	-2.77E-2
6	NO	6.45E-3	HNO2	-8.44E-3	NO	3.08E-2	O3	-9.29E-3	HNO2	-8.58E-3	NO	2.06E-2
7	SO2	2.07E-3	N2O5	1.99E-3	HNO2	-1.35E-2	N2O5	4.07E-3	N2O5	2.20E-3	HNO2	-1.29E-2
8	N2O5	1.19E-3	SO2	1.69E-3	N2O5	3.12E-3	PAN	-3.67E-3	SO2	1.78E-3	H2O2	-2.29E-3
9	PAN	-3.89E-4	H2O2	-5.08E-4	CH3O2	-7.22E-4	SO2	1.88E-3	PAN	-7.00E-4	N2O5	1.99E-3
10	CH3O2	-3.48E-4	PAN	-4.63E-4	PAN	-7.12E-4	H2O2	-5.99E-4	CH3O2	-5.96E-4	SO2	1.82E-3
11	H2O2	-2.58E-4	CH3O2	-4.53E-4	SO2	5.38E-4	CH3O2	-3.46E-4	H2O2	-4.16E-4	CH3O2	-8.32E-4
12	HNO3	1.17E-4			HNO3	1.52E-4	HNO3	1.10E-4	HNO3	1.01E-4	PAN	-3.80E-4
13					OH	-1.05E-4					OH	-1.08E-4
14											HNO3	1.01E-4

(c) Reactive hydrocarbon species

(0)												
Rank	HC	CHO	A	LD2	]	PAR	О	LE	I	SOP	CH3C	COOH
1	HCHO	1.22E+0	HCHO	-1.38E-2	HCHO	2.41E-3	HCHO	2.92E-2	HCHO	8.64E-2	НСНО	-4.60E-1
2	O3	-1.22E-2	O3	-1.96E-3	O3	3.01E-4	O3	4.67E-3	O3	9.33E-3	CH3COOH	-2.75E-1
3	HO2	-5.63E-3	HO2	-8.05E-4	HO2	1.39E-4	HO2	1.69E-3	HO2	5.00E-3	NO2	5.68E-2
4	NO2	4.40E-3	HNO2	-4.68E-4			HNO2N	9.83E-4	HNO2N	2.95E-3	O3	-5.32E-2
5	HNO2	-3.63E-3	NO2	3.66E-4			O2	-7.41E-4	O2	-2.57E-3	HO2	-2.61E-2
6	H2O2	2.21E-3	NO	2.76E-4			NO	-6.26E-4	NO	-1.52E-3	NO	1.92E-2
7	NO	5.82E-4					H2O2	1.74E-4	H2O2	5.09E-4	HNO2	-1.52E-2
8	SO2	-2.61E-4					SO2	-1.35E-4	SO2	-4.03E-4	N2O5	3.45E-3
9	CH3O2	-2.09E-4							CH3O2	1.07E-4	H2O2	-2.35E-3
10	PAN	-1.53E-4									SO2	2.18E-3
11											PAN	-1.76E-3
12											CH3O2	-7.92E-4
13											HNO3	1.23E-4
14											OH	-1.23E-4

Table 15. 1-hr average sensitivities of O<sub>3</sub> and indicators to uptake coefficients of individual species under various aerosol conditions. Only those sensitivities greater than 1.0E-4 are shown.

(a) O3

	Remote		Mari	ine	Ru	ral	Url	oan	Heavily-	polluted
Rank	Species	$\overline{S}$ kl	Species	$\overline{S}$ kl	Species	$\overline{S}_{kl}$	Species	$\overline{S}_{kl}$	Species	$\overline{S}_{kl}$
1	O3	-6.55E-4	O3	-1.31E-3	O3	-6.30E-3	O3	-4.67E-2	O3	-1.04E-1
2			H2O2	-2.76E-4	HO2	-2.13E-3	HCHO	-2.49E-2	HCHO	-6.34E-2
3			HO2	-2.54E-4	H2O2	-1.74E-3	HO2	-1.28E-2	NO2	-8.06E-3
4					HCHO	-3.59E-4	H2O2	-1.43E-3	NO	7.51E-3
5							CH3O2	-3.56E-4	HO2	-3.47E-3
6							NO	3.46E-4	N2O5	-2.15E-3
7							NO2	-2.98E-4	HNO2	-1.28E-3
8							N2O5	-2.96E-4	H2O2	-2.79E-4
9							HNO2	-2.23E-4	SO2	1.98E-4
10							SO2	2.08E-4		

(b) NOy

	Rural		Urban		Heavily-po	lluted
Rank	Species	$\overline{S}_{kl}$	Species	$\overline{S}_{kl}$	Species	$\overline{S}_{kl}$
1	HNO3	-1.31E-2	HNO3	-2.19E-2	NO2	-3.55E-2
2	NO2	-1.13E-3	NO2	-1.11E-2	HNO3	-9.09E-3
3	HO2	7.86E-4	НСНО	4.59E-3	NO	-8.92E-3
4	H2O2	6.99E-4	N2O5	-3.18E-3	HCHO	7.21E-3
5	N2O5	-6.19E-4	PAN	-3.06E-3	O3	2.89E-3
6	NO	-4.08E-4	NO	-2.83E-3	N2O5	-2.78E-3
7	PAN	-1.91E-4	HO2	2.45E-3	PAN	-1.27E-3
8	HNO2	-1.69E-4	O3	1.17E-3	HO2	4.29E-4
9	НСНО	1.50E-4	HNO2	-3.18E-4	HNO2	-2.23E-4
10			H2O2	2.92E-4		

(c) O3/NOz

	Rural		Urban		Heavily-pol	luted
Rank	Species	$\overline{S}_{kl}$	Species	$\overline{S}_{kl}$	Species	$\overline{S}_{kl}$
1	HNO3	3.72E-2	HNO3	8.18E-2	HNO3	2.22E-1
2	HO2	1.96E-2	НСНО	7.71E-2	НСНО	1.76E-1
3	H2O2	1.75E-2	HO2	4.53E-2	O3	-8.08E-2
4	O3	-4.34E-3	O3	-3.73E-2	PAN	2.96E-2
5	НСНО	2.02E-3	PAN	1.11E-2	NO2	-1.03E-2
6	SO2	-5.90E-4	H2O2	6.10E-3	HO2	1.01E-2
7	N2O5	5.40E-4	HNO2	1.22E-3	HNO2	6.96E-3
8	HNO2	3.86E-4	CH3O2	1.03E-3	NO	4.73E-3
9	CH3O2	2.71E-4	SO2	-9.12E-4	H2O2	9.61E-4
10	N2O5	2.38E-4	N2O5	8.94E-4	SO2	-9.29E-4
11	NO2	1.75E-4	NO	3.25E-4	N2O5	-2.60E-4
12	OH	1.31E-4			CH3O2	2.01E-4

(d) H2O2/HNO3

	Rural		Urban		Heavily-polluted		
Rank	Species	$\overline{S}_{kl}$	Species	$\overline{S}$ kl	Species	$\overline{S}_{kl}$	
1	H2O2	-5.48E-1	H2O2	-3.46E+0	H2O2	-2.99E+0	
2	HNO3	6.49E-2	HNO3	4.63E-1	HNO3	9.01E-1	
3	HO2	2.02E-2	НСНО	-6.19E-2	НСНО	-3.44E-1	
4	НСНО	2.06E-3	HO2	-1.64E-2	O3	-1.13E-1	
5	O3	1.99E-3	NO2	1.59E-2	NO2	8.37E-2	
6	SO2	-6.10E-4	O3	-9.60E-3	NO	3.09E-2	
7	NO2	5.40E-4	N2O5	8.05E-3	HO2	-1.75E-2	
8	N2O5	4.91E-4	NO	4.21E-3	N2O5	1.07E-2	
9	HNO2	4.39E-4	HNO2	5.31E-4	HNO2	-3.93E-3	
10	CH3O2	2.86E-4	CH3O2	-3.90E-4	SO2	-7.52E-4	
11	NO	1.93E-4	SO2	-2.80E-4	PAN	-4.97E-4	
12					CH3O2	-4.06E-4	

Appendix A-1. Gas-phase reactions in MaTChM, taken from Zaveri (1997) and references therein.

No.	Reaction	Rate constant, cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>
	Inorganic Reactions	
G1.	$NO_2 + hv \rightarrow NO + O(^3P)$	Radiation Dependent
G2.	$O(^{3}P) \xrightarrow{\circ 2,M} O_{3}$	1.4E+03 exp(1175/T)
G3.	$O_3 + NO \rightarrow NO_2$	1.8E-12exp(-1370/T)
G4.	$o(^3P) + No_2 \rightarrow No$	9.3E-12
G5.	$O(^3P) + NO_2 \xrightarrow{M} NO_3$	1.6E-13 exp(687/T)
G6.	$O(^{3}P)+ NO \xrightarrow{M} NO_{2}$	2.2E-13 exp(602/T)
G7.	$O_3 + NO_2 \rightarrow NO_3$	1.2E-13exp(-2450/T)
G8.	$O_3 + hv \rightarrow O(^3P)$	Radiation Dependent
G9.	$O_3 + hv \rightarrow O(^1D)$	Radiation Dependent
G10.	$O(1D) \xrightarrow{M} O(^{3}P)$	1.19E+08 exp(390/T)
G11.	$O(^{1}D) + H_{2}O \rightarrow 2OH$	2.2E-10
G12.	$O_3 + OH \rightarrow HO_2$	1.6E-12 exp(-940/T)
G13.	$O_3 + HO_2 \rightarrow OH$	1.4E-14 exp(-580/T)
G14.	$NO_3 + hv \rightarrow 89NO_2 + .89O + .11NO$	Radiation Dependent
G15.	$NO_3 + NO \rightarrow 2NO_2$	1.3E-11 exp(250/T)
G16.	$NO_3 + NO_2 \rightarrow NO + NO_2$	2.5E-14 exp(-1230/T)
G17.	$NO_3 + NO_2 \xrightarrow{M} N_2O_5$	5.3E-13 exp(256/T)
G18.	$N_2O_5 + H_2O \rightarrow 2HNO_3$	1.3E-21
G19.	$N_2O_5 \xrightarrow{M} NO_3 + NO_2$	3.5E+14exp(-10897/T)
G20.	$NO + NO \xrightarrow{o 2} 2NO_2$	1.8E-20 exp(530/T)
G21.	$NO + NO_2 + H_2O \rightarrow 2HONO$	4.4E-40
G22.	$OH + NO \xrightarrow{M} HONO$	4.5E-13 exp(806/T)
G23.	$HONO + hv \rightarrow OH + NO$	Radiation Dependent
G24.	$OH + HONO \rightarrow NO_2$ $HONO + HONO \rightarrow NO + NO$	6.6E-12
G25.	$HONO + HONO \rightarrow NO + NO_2$ $OH + NO_2 \xrightarrow{M} HNO_3$	1.0E-20
G26.	$OH + NO_2 \xrightarrow{M} NO_3$ $OH + HNO_3 \xrightarrow{M} NO_3$	1.0E-12 exp(713/T)
G27. G28.	$OH + HNO_3 \longrightarrow NO_3$ $HO_2 + NO \rightarrow OH + NO_2$	5.1E-15 exp(1000/T) 3.7E-12 exp(240/T)
G29.	$HO_2 + NO_2 \xrightarrow{M} PNA$	1.2E-13 exp(240/T)
	$PNA \xrightarrow{M} HO_2 + NO_2$	• • • •
G30.		4.8E+13exp(-10121/T)
G31. G32.	$OH + PNA \rightarrow NO_2$ $HO_2 + HO_2 \rightarrow H_2O_2$	1.3E-12 exp(380/T) 5.9E-14 exp(1150/T)
G32.	$HO_2 + HO_2 \rightarrow H_2O_2$ $HO_2 + HO_2 + H_2O \rightarrow H_2O_2$	2 .2E-38 exp(5800/T)
G33.	$H_2O_2 + hv \rightarrow 2OH$	Radiation Dependent
G35.	$OH + H2O2 \rightarrow HO2$	3.1E-12 exp(-187/T)
G36.	$CO + OH \xrightarrow{a^2} HO_2$	2.2E-13
	Formaldehyde Reactions	
G37.	FORM + OH $\xrightarrow{o^2}$ HO <sub>2</sub> + CO	1.0E-11
G38.	$FORM + hv \xrightarrow{2\sigma^2} 2HO_2 + CO$	Radiation Dependent
G39.	$FORM + hv \rightarrow CO$	Radiation Dependent
G40.	$FORM + O(^{3}P) \rightarrow OH + HO_{2} + CO$	3.0E-11 exp(-1550/T)
G41.	$FORM + NO_3 \xrightarrow{o 2} HNO_3 + HO_2 + CO$	6.3E-16

Read 1.4E+03 exp(1175/T) as 1.4 x  $10^3$  e<sup>1175/T</sup>.

No.	Reaction	Rate constant, cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>
	Higher Molecular Weight Aldehyde Reactions	
G42.	$ALD2 + O(^{3}P) \xrightarrow{o 2} C2O3 + OH$	1.2E-11 exp(-986/T)
G43.	$ALD2 + OH \rightarrow C2O3$	7.0E-12 exp(250/T)
G44.	$ALD2 + NO_3 \xrightarrow{\qquad o \ 2} C2O3 + HNO_3$	2.5E-15
G45.	$ALD2 + hv \xrightarrow{2 \circ 2} CH_3O_2 + HO_2 + CO$	Radiation dependent
G46.	$C2O3 + NO \xrightarrow{\circ 2} CH_3O_2 + NO_2$	5.4E-12 exp(250/T)
G47.	$C2O3 + NO_2 \rightarrow PAN$	8.0E-20 exp(5500/T)
G48.	$PAN \rightarrow C2O3 + NO_2$	9.4E+16exp(-14000/T)
G49.	$C2O3 + C2O3 \rightarrow 2CH_3O_2 + O_2$	2.0E-12
G50.	$C2O3 + HO_2 \rightarrow .79CH_3O_2 + .79OH$	6.5E-12
	Alkane Reactions	
G51.	$C2H6 + OH \rightarrow ETHP$	$T^2$ 1.37E-17exp(-444/T)
G52.	$PAR + OH \rightarrow .87XO2 + .13XO2N + .11HO_2 + .11ALD2 + .76ROR11PAR$	8.1E-13
G53.	$ROR \rightarrow 1.1ALD2 + .96XO2 + .94HO_2 + .04XO2N + .02ROR - 2.1PAR$	1.0E+15 exp(-8000/T)
G54.	$ROR \rightarrow HO_2$	1.6E+3
G55.	$ROR + NO_2 \rightarrow NITRATES$	1.5E-11
	Alkene Reactions	
G56.	OLE+ $O(^{3}P) \rightarrow .63ALD2 + .38HO_{2} + .28XO2 + .3CO + .2HCHO + .02XO2N + .22PAR + .2OH$	1.2E-11 exp(-324/T)
G57.	OLE + OH $\rightarrow$ CH <sub>3</sub> O <sub>2</sub> + ALD2 - PAR	5.2E-12 exp(504/T)
G58.	OLE + $O_3 \rightarrow .5ALD2 + .524HCHO + .33CO + .228HO_2 + .10H + .216CH_3O_2 - PAR$	1.4E-14 exp(-2105/T)
G59.	OLE + $NO_3 \rightarrow 91XO2 + HCHO + ALD2 + .09XO2N + NO_2 - PAR$	7.7E-15
G60.	ETH + $O(^{3}P) \rightarrow .3HCHO + .7CH_{3}O_{2} + CO + HO_{2} + .3OH$	1.0E-11 exp(-792/T)
G61.	ETH + OH $\rightarrow$ XO2 + 1.56HCHO + HO <sub>2</sub> + .22ALD2	2.0E-12 exp(411/T)
G62.	$ETH + O_3 \rightarrow HCHO + .42CO + .12HO_2$	1.3E-14 exp(-2633/T)
G63.	Aromatic Reactions $TOL + OH \rightarrow .08XO2 + .36CRES + .44HO_2 + .56TO2$	2.1E-12 exp(322/T)
G64.	TO2 + NO $\rightarrow$ .9NO2 + .9OPEN + .9HO2	8.1E-12
G65.	$TO2 \rightarrow CRES$	4.2
G66.	$CRES + OH \rightarrow .4CRO + .6XO2 + .6HO2 + .3OPEN$	4.1E-11
G67.	$CRES + NO_3 \rightarrow CRO + HNO_3$	2.2E-11
G68.	$CRO + NO_2 \rightarrow NITRATES$	1.4E-11
G69. G70.	$XYL + OH \rightarrow .7OH + .5XO2 + .2CRES + .8MGLY + 1.1PAR + .3TO2$ $OPEN + OH \rightarrow XO2 + C2O3 + 2HO_2 + 2CO + HCHO$	1.7E-11 exp(116/T) 3.0E-11
G70.	$OPEN + hv \rightarrow C2O3 + CO + HO2$	Radiation dependent
G72.	OPEN+ O <sub>3</sub> →.03ALD2+.62C2O3+.7HCHO+.03XO2 +.69CO+.08OH+.76HO <sub>2</sub> +.2MGLY	5.4E-17exp(-500/T)
G73.	$MGLY + OH \rightarrow XO2 + C2O3$	1.7E-11
G74.	$MGLY + hv \rightarrow C2O3 + CO + HO_2$	Radiation dependent
	Isoprene Reactions	
G75.	$ISOP + O(^{3}P) \rightarrow .6HO_{2} + .8ALD2 + .5SOLE + .5XO2 + .5CO + .45ETH + .9PAR$	1.8E-11
G76.	$ISOP + OH \rightarrow \!\! HCHO \!\! + \!\! XO2 \!\! + \!\! .67HO_2 \!\! + \!\! .4MGLY \!\! + \!\! .2C2O3 + \!\! ETH \!\! + \!\! .2ALD2 \!\! + \!\! .13XO2N$	9.6E-11
G77.	$ISOP + O_3 \rightarrow HCHO + .4ALD2 + .55ETH + .2MGLY + .06CO + .1PAR + .44HO_2 + .1OH$	1.2E-17
G78.	$ISOP + NO_3 \rightarrow XO2N + NITRATES$	3.2E-13
670	Operator Reactions VO2 - NO - NO	9.1E.12
G79.	$XO2 + NO \rightarrow NO_2$ $YO2 + YO2 \rightarrow Products$	8.1E-12
G80. G81.	$XO2 + XO2 \rightarrow Products$ $XO2N + NO \rightarrow NITRATES$	1.7E-14 exp(1300/T) 6.8E-13
501.	AACMAT TATO TATABATAN	0.0E 10

No.	Reaction	Rate constant, cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>
	Condensed DMS Chemistry	
G82.	$DMS + OH \rightarrow CH_3SCH_2OO + H_2O$	9.64E-12 exp(-234/T)
G83.	$DMS + NO_3 \rightarrow CH_3SCH_2OO + HNO_3$	1.40E-13 exp(500/T)
G84.	$DMS + O(^{3}P) \rightarrow CH_{3}SO_{2} + CH_{3}O_{2}$	.26E-11 exp(409/T)
G85.	$DMS + OH \rightarrow aCH_3SO_2 + aCH_3O_2 + (1\text{-}a)DMSO + (1\text{-}a)HO_2$	1.7E-12 (see note)
G86.	$CH_3SCH_2OO + NO \rightarrow CH_3SO_2 + HCHO + NO_2$	8.0E-12
G87.	$CH_3SCH_2OO + CH_3O_2 \rightarrow CH_3SO_2 + CH_3O + HCHO$	1.8E-13
G88.	$CH_3SCH_2OO + HO_2 \rightarrow CH_3SCH_2OOH$	1.5E-12
G89.	$DMSO + OH \rightarrow bCH_3 SO_2 H + bCH_3 O_2 + (1-b) DMSO_2 + (1-b) HO_2$	5.8E-11 (see note)
G90.	$DMSO_2 + OH \rightarrow CH_3S(O)_2CH_2OO + H_2O$	1.0E-14
G91.	$CH_3S(O)_2CH_2OO + NO \rightarrow CH_3SO_2 + HCHO + NO_2$	5.0E-12
G92.	$\mathrm{CH_{3}S(O)_{2}CH_{2}OO} + \mathrm{CH_{3}O_{2}} \rightarrow \mathrm{CH_{3}SO_{2}} + \mathrm{HCHO} + \mathrm{CH_{3}O}$	1.8E-13
G93.	$CH_3S(O)_2CH_2OO+HO_2\rightarrow CH_3S(O)_2CH_2OOH+O_2$	1.5E-12
G94.	$CH_3SO_2H + HO_2 \rightarrow CH_3SO_2 + H_2O_2$	1.0E-15
G95.	$\text{CH}_3\text{SO}_2\text{H} + \text{NO}_3 \rightarrow \text{CH}_3\text{SO}_2 + \text{HNO}_3$	1.0E-13
G96.	$CH_3SO_2H + CH_3O_2 \rightarrow CH_3SO_2 + CH_3OOH$	1.0E-15
G97.	$CH_3SO_2H + OH \rightarrow CH_3SO_2 + H_2O$	1.6E-11
G98.	$\text{CH}_3\text{SO}_2\text{H} + \text{CH}_3\text{SO}_3 \rightarrow \text{CH}_3\text{SO}_2 + \text{MSA}$	1.0E-13
G99.	$CH_3SO_2 \xrightarrow{M} SO_2 + CH_3O_2$	4.53E+13exp(-8656/T)
G100.	$CH_3SO_2 + NO_2 \rightarrow CH_3SO_3 + NO$	1.0E-14
G101.	$CH_3SO_2 + O_3 \rightarrow CH_3SO_3 + O_2$	1.0E-15
G102.	$CH_3SO_2 + HO_2 \rightarrow CH_3SO_3 + OH$	2.5E-13
G103.	CH <sub>3</sub> SO <sub>2</sub> + CH <sub>3</sub> O <sub>2</sub> →CH <sub>3</sub> SO <sub>3</sub> +CH <sub>3</sub> O+ O <sub>2</sub>	2.5E-13
G104.	$CH_3SO_2 + OH \rightarrow MSA$	5.0E-11
G105.	$CH_3SO_2 + O_2 \xrightarrow{M} CH_3S(O)_2OO$	2.6E-18
G106.	$CH_3S(O)_2OO \xrightarrow{M} CH_3SO_2 + O_2$	3.3
G107.	$CH_3S(O)_2OO + NO \rightarrow CH_3SO_3 + NO_2$	1.0E-11
G108.	CH <sub>3</sub> S(O) <sub>2</sub> OO+CH <sub>3</sub> O <sub>2</sub> →CH <sub>3</sub> SO <sub>3</sub> +CH <sub>3</sub> O+O <sub>2</sub>	5.5E-12
G109.	$CH_3S(O)_2OO+HO_2\rightarrow CH_3S(O)_2OOH+O_2$	2.0E-12
G110.	$CH_3SO_3 \xrightarrow{M} H_2SO_4 + CH_3O_2$	1.6E-1
G111.	$CH_3SO_3 + NO_2 \rightarrow MSA + HNO_3$	3.0E-15
G112.	$CH_3SO_3 + NO \rightarrow MSA + HNO_2$	3.0E-15
G113.	$CH_3SO_3 + HO_2 \rightarrow MSA + O_2$	5.0E-11
G114.	$CH_3SO_3 + HCHO \xrightarrow{\circ 2} MSA + HO_2 + CO$	1.6E-15
G115.	$SO_2 + OH \xrightarrow{M} H_2SO_4 + HO_2$	Troe expression
	Methane and Methylperoxyl Radical Reactions	
G116.	$CH_4 + OH \xrightarrow{\sigma^2} CH_3O_2$	$T^2$ 6.95E-18exp(-1280/T)
G117.	$CH_3O_2 + NO \rightarrow HCHO + HO_2 + NO_2$	4.2E-12 exp(180/T)
G118.	$CH_3O_2 + HO_2 \rightarrow CH_3OOH$	7.7E-14 exp(1300/T)
G119.	$CH3O2 + CH3O2 \rightarrow 1.5HCHO + HO2$	1.9E-13 exp(220/T)
G120.	$CH_3O_2 + C2O_3 \rightarrow HCHO + .5HO_2 + .5CH_3O_2 + CH_3OOH$	9.6E-13 exp(220/T)
G121.	$CH_3OOH + hv \rightarrow HCHO + HO_2 + OH$	5.57E-06
G122.	CH <sub>3</sub> OOH+OH→.5CH <sub>3</sub> O <sub>2</sub> +.5HCHO+.5OH	1.0E-11
G123.	$ETHP + NO \rightarrow ALD2 + HO_2 + NO_2$	4.2E-12 exp(180/T)
G124.	$ETHP + HO_2 \rightarrow Products$	7.7E-14 exp(1300/T)
G125.	ETHP + C2O3 $\rightarrow$ ALD2 + .5HO <sub>2</sub> + .5CH <sub>3</sub> O <sub>2</sub> + .5CH <sub>3</sub> COOH	3.4E-13 exp(220/T)

Note:  $a = 5E + 5/(5E + 5 + [O_2] \times 3E - 12)$ ;  $b = 1.5E + 7/(1.5E + 7 + [O_2] \times 1.2E - 12)$ 

Appendix A-2. Aqueous-phase reactions in MaTChM, taken from Zaveri (1997) and references therein.

No.	Reaction	Rate Constant, M <sup>n</sup> s <sup>-1</sup>
L1.	Oxygen-Hydrogen Chemistry $H_2O_2 + hn \rightarrow 2OH$	5.4E-7
L2.	$O_3 + hn \xrightarrow{H_2O} H_2O_2 + O_2$	1.2E-5
L3.	$OH + HO_2 \rightarrow H_2O + O_2$	1.07E+12 exp(-1500/T)
L4.	$OH + O_2^- \rightarrow OH^- + O_2$	1.53E+12 exp(-1500/T)
L5.	$OH + H2O2 \rightarrow H2O + HO2$ $HO2 + HO2 \rightarrow H2O + O3$	8.12E+9 exp(-1700/T)
L6.	$HO_2 + HO_2 \rightarrow H_2O_2 + O_2$ $H_2O$	2.40E+9 exp(-2365/T)
L7.	$\text{HO}_2 + \text{O}_2^- \xrightarrow{H^2O} \text{H}_2\text{O}_2 + \text{O}_2 + \text{OH}^-$	1.53E+10 exp(-1500/T)
L8.	$O_2^- + O_2^- \xrightarrow{2H_2O} H_2O_2 + O_2 + 2OH^-$	3.0E-1
L9.	$HO_2 + H_2O_2 \rightarrow OH + O_2 + H_2O$	5.0E-1
.10.	$O_2^- + H_2O_2 \rightarrow OH + O_2 + OH^-$	1.3E-1
.11.	$OH + O_3 \rightarrow HO_2 + O_2$	2.0E+9
.12.	$HO_2 + O_3 \rightarrow OH + 2O_2$	5.0E+3
.13.	$O_2^- + O_3 \rightarrow OH + 2O_2 + OH^-$	2.30E+11exp(-1500/T)
.14.	$OH^- + O_3 \xrightarrow{H^2O} O_2^- + HO_2$	7.0E+1
ـ15.	$HO_2^- + O_3 \rightarrow OH + O_2^- + O_2$	2.8E+6
.16.	$H_2O_2 + O_3 \rightarrow H_2O + 2O_2$	7.8E-3 [O <sub>3</sub> ] <sup>-0.5</sup>
	<u>Carbonate Chemistry</u>	
<b>L</b> 17.	$HCO_3^- + OH \rightarrow H_2O + CO_3^-$	9.11E+9 exp(-1910/T)
.18.	$HCO_3^- + O_2^- \rightarrow HO_2^- + CO_3^-$	1.5E+6
<b>.</b> 19.	$CO_3^- + O_2^- \xrightarrow{H2O} HCO_3^- + O_2 + OH^-$	6.14E+10 exp(-1500/T)
L20.	$CO_3^- + H_2O_2 \rightarrow HO_2 + HCO_3^-$	1.03E+10exp(-2820/T)
	Chlorine Chemistry	
L21.	$Cl^- + OH \rightarrow ClOH^-$	6.6E+11 exp(-1500/T)
L22.	$CIOH^- \rightarrow CI^- + OH$	6.1E+9
L23.	$CIOH^- \xrightarrow{H^+} CI + H_2O$	2.1E+10 [H <sup>+</sup> ]
L24.	$CI \xrightarrow{H_2O} CIOH^- + H^+$	1.3E+3
.24. .25.	$HO_2 + Cl_2^- \rightarrow 2Cl^- + O_2 + H^+$	6.91E+11 exp(-1500/T)
26.	$O_2^- + Cl_2^- \rightarrow 2Cl^- + O_2$	1.53E+11 exp(-1500/T)
.20. .27.	$O_2 + CI_2 \rightarrow 2CI + O_2$ $HO_2 + CI \rightarrow CI^- + O_2 + H^+$	• • • • • • • • • • • • • • • • • • • •
	2 2	4.76E+11 exp(-1500/T)
.28.	$H_2O_2 + Cl_2^- \rightarrow 2Cl^- + HO_2 + H^+$	1.14E+10 exp(-3370/T)
.29.	$H_2O_2 + CI \rightarrow CI^- + HO_2 + H^+$	4.5E+7
.30.	$OH^- + Cl_2^- \rightarrow 2Cl^- + OH$	$1.02E+10 \exp(-2160/T)$
	Nitrite and Nitrate Chemistry	
.31.	$NO + NO_2 \xrightarrow{H_2O} 2NO_2^- + 2H^+$	3.07E+10 exp(-1500/T)
.32.	$NO_2 + NO_2 \xrightarrow{H^2O} NO_2^- + NO_3^- + 2H^+$	1.53E+10 exp(-1500/T)
<b>.</b> 33.	$NO + OH \rightarrow NO_2^- + H^+$	3.07E+12 exp(-1500/T)
.34.	$NO2 + OH \rightarrow NO_3^- + H^+$	2.0E+11 exp(-1500/T)
.35.	$HNO_2 + hn \rightarrow NO + OH$	3.7E-5
.36.	$NO_2^- + hn \xrightarrow{H_2O} NO + OH + OH^-$	6.3E-6
<b>.</b> 37.	$HNO_2 + OH \rightarrow NO_2 + H_2O$	1.53E+11 exp(-1500/T)
.38.	$NO_2^- + OH \rightarrow NO_2 + OH^-$	1.53E+12 exp(-1500/T)
<b>.</b> 39.	$HNO_2 + H_2O_2 \xrightarrow{H^+} NO_3^- + 2H^+ + H_2O$	3.66E+13 exp(-6700/T) [H <sup>+</sup> ]
.40.	$NO_2^- + O_3 \rightarrow NO_3^- + O_2$	6.72E+15 exp(-6950/T)
<b>.</b> 41.	$NO_2^- + CO_3^- \rightarrow NO_2 + CO_3^{2-}$	4.0E+5

No.	Reaction	Rate constant, M <sup>n</sup> s <sup>-1</sup>
L42.	$NO_2^- + Cl_2^- \rightarrow NO_2 + 2Cl^-$	3.84E+10 exp(-1500/T)
L43.	$NO_2^- + NO_3 \rightarrow NO_2 + NO_3^-$	1.84E+11 exp(-1500/T)
L44.	$NO_3^- + hn \xrightarrow{H2O} NO_2 + OH + OH^-$	1.8E-7
L45.	$NO_3 + hn \rightarrow NO + O_2$	0.0
L46.	$NO_3 + HO_2 \rightarrow NO_3^- + H^+ + O_2$	6.90E+11 exp(-1500/T)
L47.	$NO_3 + O_2^- \rightarrow NO_3^- + O_2$	1.53E+11 exp(-1500/T)
L48.	$NO_3 + H_2O_2 \rightarrow NO_3^- + HO_2 + H^+$	1.20E+10 exp(-2800/T)
L49.	$NO_3 + Cl^- \rightarrow NO_3^- + Cl$	1.53E+10 exp(-1500/T)
	Methane Oxidation Chain	
L50.	$CH_2(OH)_2 + OH \xrightarrow{\circ 2} HCOOH + HO_2 + H_2O$	3.10E+11 exp(-1500/T)
L51.	$CH_2(OH)_2 + O_3 \rightarrow Products$	1.0E-1
L52.	$\text{HCOOH} + \text{OH} \xrightarrow{ o  2 } \text{CO}_2 + \text{HO}_2 + \text{H}_2\text{O}$	2.45E+10 exp(-1500/T)
L53.	$HCOOH + H_2O_2 \rightarrow Product + H_2O$	1.63E+2 exp(-5180/T)
L54.	$\text{HCOOH} + \text{NO}_3 \xrightarrow{\qquad o  2} \text{NO}_3^- + \text{H}^+ + \text{CO}_2 + \text{HO}_2$	9.68E+9 exp(-3200/T)
L55.	$HCOOH + O_3 \rightarrow CO_2 + HO_2 + OH$	5.0
L56.	$\text{HCOOH} + \text{Cl}_2^- \xrightarrow{\sigma^2} \text{CO}_2 + \text{HO}_2 + 2\text{Cl}^- + \text{H}^+$	1.24E+10 exp(-4300/T)
L57.	$\text{HCOO}^- + \text{OH} \xrightarrow{\circ 2} \text{CO}_2 + \text{HO}_2 + \text{OH}^-$	3.84E+11 exp(-1500/T)
L58.	$HCOO- + O_3 \rightarrow CO_2 + OH + O_2^-$	1.0E+2
L59.	$\text{HCOO}^- + \text{NO}_3 \xrightarrow{ o^2 } \text{NO}_3^- + \text{CO}_2 + \text{HO}_2$	9.21E+9 exp(-1500/T)
L60.	$\text{HCOO}^{\text{-}} + \text{CO}_3^{\text{-}} \xrightarrow{ o \text{ 2,H2o} } \text{CO}_2 + \text{HCO}_3^{\text{-}} + \text{HO}_2 + \text{OH}^{\text{-}}$	9.92E+9 exp(-3400/T)
L61.	$HCOO^- + Cl_2^- \xrightarrow{\sigma^2} CO_2 + HO_2 + 2Cl^-$	1.17E+10 exp(-2600/T)
L62.	$CH_3C(O)O_2NO_2 \rightarrow NO_3^- + Products$	4.0E-4
L63.	$CH_3O_2 + HO_2 \rightarrow CH_3OOH + O_2$	1.01E+10 exp(-3000/T)
L64.	$CH3O2 + O2- \xrightarrow{H2o} CH3OOH + O2 + OH-$	1.07E+10 exp(-1600/T)
L65.	$CH_3OOH + hn \xrightarrow{o^2} HCHO + OH + HO_2$	5.40E-7
L66.	$CH_3OOH + OH \rightarrow CH_3O_2 + H_2O$	8.11E+9 exp(-1700/T)
L67.	$CH_3OH + OH \rightarrow HCHO + HO_2 + H_2O$	6.91E+8 exp(-1500/T)
L68.	$CH_3OH + CO_3^- \xrightarrow{g_2} HCHO + HO_2 + HCO_3^-$	9.40E+9 exp(-4500/T)
L69.	$CH_3OH + Cl_2^{-} \xrightarrow{\sigma^2} HCHO + HO_2 + H^+ + 2Cl^-$	9.04E+9 exp(-4400/T)
L70.	$CH_3OOH + OH \rightarrow HCHO + OH + H_2O$	7.98E+9 exp(-1800/T)
L71.	$CH_3OH + NO_3 \xrightarrow{\sigma^2} NO_3^- + H^+ + HCHO + HO_2$	1.20E+10 exp(-2800/T)
	Sulfur Chemistry	
L72.	$H_2SO_3 + O_3 \rightarrow S(VI)$	2.4E+4
L73.	$HSO_3^- + O_3 \rightarrow S(VI)$	4.24E+13 exp(-5530/T)
L74.	$SO_3^{2-} + O_3 \rightarrow S(VI)$	7.43E+16exp(-5280/T)
L75.	$HSO_3^- + H_2O_2 \xrightarrow{H^+} S(VI) + H_2O$	2.9E+10 [H+] exp(-1950/T) (1 + 16[H+])
L76	$HSO_3^- + H_2O_2 \xrightarrow{HX} S(VI) + H_2O$	3.53E+7 [HCOOH] <sub>1</sub> exp(-1720/T)
	5 2 2 \ / 2	1.91E+8 [CH3COOH] <sub>l</sub> exp(-2620/T)
L77.	$S(IV) \xrightarrow{o 2, Mn^{2+}, Fe^{3+}} S(VI)$	pH 0-5, 2.6E+3 [Fe <sup>3+</sup> ] + 7.5E+
		$2 [Mn^{2+}] + 1.0E + 10 [Fe^{3+}][Mn^{2+}]$
1.70	$SO_3^{2-} + OH \xrightarrow{\sigma^2} SO_5^- + OH^-$	pH >5, 7.5E+2 [Mn2+] + 2E10 [Fe <sup>3+</sup> ][Mn <sup>2+</sup> ]
L78.	-	7.06E+11 exp(-1500/T)
L79.	$HSO_3^- + OH \xrightarrow{\sigma^2} SO_5^- + H_2O$	6.44E+11 exp(-1500/T)
L80.	$SO_5^- + HSO_3^- \xrightarrow{\sigma^2} HSO_5^- + SO_5^-$	9.88E+9 exp(-3100/T)

No.	Reaction	Rate constant, M <sup>n</sup> s <sup>-1</sup>
L81.	$SO_5^- + SO_3^{2-} \xrightarrow{\qquad o \ 2} HSO_5^- + SO_5^-$	1.07E+10 exp(-2000/T)
L82.	$SO_5^- + O_2^- \xrightarrow{H^{2o}} HSO_5^- + OH^- + O_2$	1.53E+10 exp(-1500/T)
L83.	$SO_5^- + HCOOH \xrightarrow{\qquad o \ 2} HSO_5^- + CO_2 + HO_2$	1.06E+10 exp(-5300/T)
L84.	$SO_5^- + HCOO^- \xrightarrow{\sigma^2} HSO_5^- + CO_2 + O_2^-$	9.45E+9 exp(-4000/T)
L85.	$SO_5^- + SO_5^- \to 2SO_4^- + O_2$	3.07E+10 exp(-1500/T)
L86.	$HSO_5^- + HSO_3^- \xrightarrow{H^+} 2SO_4^{2-} + 3H^+$	6.27E+14 exp(-4750/T)
L87.	$HSO_5^- + OH \rightarrow SO_5^- + H_2O$	9.99E+9 exp(-1900/T)
L88.	$HSO_5^- + SO_4^- \to SO_5^- + SO_4^{2-} + H^+$	5.0E+4
L89.	$HSO_5^- + NO_2^- \rightarrow HSO_4^- + NO_3^-$	1.52E+9 exp(-6650/T)
L90.	$HSO_5^- + Cl^- \rightarrow SO_4^{2-} + Product$	3.38E+7 exp(-7050/T)
L91.	$SO_4^- + HSO_3^- \xrightarrow{-0.2} SO_4^{2-} + H^+ + SO_5^-$	2.00E+11 exp(-1500/T)
L92.	$SO_4^- + SO_3^{2-} \xrightarrow{\sigma^2} SO_4^{2-} + SO_5^-$	8.13E+10 exp(-1500/T)
L93.	$SO_4^- + HO_2 \rightarrow SO_4^{2-} + H^+ + O_2$	7.67E+11 exp(-1500/T)
L94.	$SO_4^- + O_2^- \to SO_4^{2-} + O_2$	7.67E+11 exp(-1500/T)
L95.	$SO_4^- + OH^- \rightarrow SO_4^{2-} + OH$	1.23E+10 exp(-1500/T)
L96.	$SO_4^- + H_2O_2 \rightarrow SO_4^{2-} + H^+ + HO_2$	9.86E+9 exp(-2000/T)
L97.	$SO_4^- + NO_2^- \rightarrow SO_4^{2-} + NO_2$	1.35E+11 exp(-1500/T)
L98.	$SO_4^- + HCO_3^- \rightarrow SO_4^{2-} + H^+ + CO_3^-$	1.04E+10 exp(-2100/T)
L99.	$SO_4^- + HCOO^- \xrightarrow{\sigma^2} SO_4^{2-} + CO_2 + HO_2$	2.61E+10 exp(-1500/T)
L100.	$SO_4^- + Cl^- \rightarrow SO_4^{2-} + Cl$	3.07E+10 exp(-1500/T)
L101.	$SO_4^- + HCOOH \xrightarrow{\circ 2} SO_4^{2-} + H^+ + CO_2 + HO_2$	1.21E+10 exp(-2700/T)
L102.	$S(IV) + CH_3C(O)O_2NO_2 \rightarrow S(VI)$	6.7E-3 see note [a]
L103.	$HSO_3^- + CH_3OOH \xrightarrow{H^+} SO_4^{2^-} + 2H^+ + Products$	6.56E+12 exp(-3800/T)
L104.	$HSO_3^- + CH_3C(O)OOH \rightarrow SO_4^{2-} + H^+ + Products$	3.37E+13 exp(-4000/T) see note [a]
L105.	$S(IV) + HO_2 \rightarrow S(VI) + OH$	1.0E+6
L106.	$S(IV) + O_2^- \xrightarrow{H^{2o}} S(VI) + OH + OH^-$	1.0E+5
L107.	$SO4^- + CH_3OH \xrightarrow{\circ 2} SO_4^{2-} + HCHO + H^+ + HO_2$	1.05E+10 exp(-1800/T)
L108.	$2HSO_3^- + NO_3 \xrightarrow{o 2} NO_3^- + 2H^+ + SO_4^{2-} + SO_4^-$	1.0E+8
L109.	$2NO_2 + HSO_3^- \xrightarrow{H2o} SO_4^{2-} + 3H^+ + 2NO_2^-$	2.0E+6
L110a.	$S(IV) + N(III) \rightarrow S(VI) +$	$pH \le 3  1.42E+2 [H^+]^{0.5}$
L110b.	$2HSO_3^- + NO_2^- \rightarrow OH^- + HDS$	pH > 3 4.76E+3 [H <sup>+</sup> ] exp(-6100/T)
L111.	$HCHO + HSO_3^- \rightarrow HOCH_2SO_3^-$	4.01E+9 exp(-4900/T)
L112	$\text{HCHO} + \text{SO}_3^{2^-} \xrightarrow{H^{2o}} \text{HOCH}_2\text{SO}_3^- + \text{OH}^-$	1.05E+10 exp(-1800/T)
L113.	$HOCH_2SO_3^- + OH- \rightarrow SO_3^{2-} + HCHO + H_2O$	1.30E+10 exp(-4500/T)
L114.	$\text{HOCH}_2\text{SO}_3^- + \text{OH} \xrightarrow{ \sigma \text{ 2} } \text{SO}_5^- + \text{HCHO} + \text{H}_2\text{O}$	2.15E+11 exp(-1500/T)
L115.	$HSO_3^- + Cl_2^- \xrightarrow{o 2} SO_5^- + 2Cl^- + H^+$	5.22E+10 exp(-1500/T)
L116.	$SO_3^{2^-} + Cl_2^- \xrightarrow{\sigma^2} SO_5^- + 2Cl^-$	2.45E+10 exp(-1500/T)
L117.	$DMS + OH \rightarrow 0.5DMSO + 0.5DMS + 0.5H_2O$	1.9E+10
L118.	$DMS + H_2O_2 \rightarrow DMSO + H_2O$	3.5E-2
L119.	$DMS + O_3 \rightarrow DMSO + O_2$ $DMSO + O_3 \rightarrow DMSO + O_3$	6.72E+15 exp(-4832/T)
L120.	$DMSO + O_3 \rightarrow DMSO2 + O_2$	5.7

a For non-elementary rate expression,  $[S(IV)] = [SO_2]_W$ ;  $[S(VI)] = [H_2SO_4]_W$ ;  $[N(III)] = [HNO_2]_W$ .

Appendix A-3. Dissolution equilibria in MaTChM, taken from Zaveri (1997) and references therein.

No.	Dissolution Equilibria	Henry's Law Constant, H $[M_{(1)}]/[M_{(g)}]$
E1.	$HNO_3(g) \iff HNO_3(l)$	5.1E+6 exp(8700FT)
E2.	$HNO_2(g) \iff HNO_2(l)$	1.2E+3 exp(4780FT)
E3.	$SO_2(g) \iff H_2SO_3$	3.0E+1 exp(3120FT)
E4.	$H_2SO_4(g) \iff H_2SO_4(l)$	1.0E+3
E5.	$HCOOH(g) \iff HCOOH(l)$	9.04E+4 exp(5700FT)
E6.	$CH_3COOH(g) \iff CH_3COOH(l)$	1.0E+3 (assumed)
E7.	$CO_2(g) \iff H_2CO_3$	8.31E-1 exp(2420FT)
E8.	$MSA(g) \iff MSA(l)$	1.0E+3 (assumed)
E9.	$NH_3(g) \iff NH_4OH$	1.38E+3 exp(4181FT)
E10.	$HO_2(g) \iff HO_2(l)$	4.89E+4 exp(6640FT)
E11.	$H_2O_2(g) \iff H_2O_2(l)$	2.44E+1 exp(7514/T - 13.63)
E12.	$HCHO(g) \iff HCHO(l)$	1.54E+5 exp(6460FT)
E13.	$NO_2(g) \iff NO_2(1)$	2.44E-1 exp(2500FT)
E14.	$NO(g) \iff NO(1)$	4.64E-2 exp(1480FT)
E15.	$NO_3(g) \iff NO_3(1)$	3.67E+2
E16.	$PAN(g) \iff PAN(l)$	7.1E+1 exp(5910FT)
E17.	$O_3(g) \iff O_3(1)$	2.76E-1 exp(2300FT)
E18.	$OH(g) \iff OH(1)$	6.10E+2 exp(5280FT)
E19.	$CH_3O_2(g) \iff CH_3O_2(l)$	1.46E+2 exp(5600FT)
E20.	$CH_3OH(g) \iff CH_3OH(l)$	5.38E+3 exp(4900FT)
E21.	$DMS(g) \iff DMS(l)$	1.17E-4 exp(3480/T)
E22.	$DMSO(g) \iff DMSO(l)$	2.44E+7
E23.	$DMSO2(g) \iff DMSO2(1)$	1.0E+4 (assumed)
E24.	$CH_3OOH(g) \iff CH_3OOH(1)$	5.55E+3 exp(5610FT)
E25.	$CH_3C(O)OOH(g) \iff CH_3C(O)OOH(l)$	1.15E+4 exp(6170FT)
E26.	$N_2O_5(g) \rightarrow 2HNO_3(w)$	1.0E+4 (assumed)
E27.	$NTR(g) \iff NTR(l)$	1.0 (assumed)
E28.	$HCl(g) \iff HCl(l)$	1.77E+4 exp(2020FT)
E29.	$Cl(g) \iff Cl(l)$	1.77E+4 exp(2020FT)

Appendix A-4. Dissociation equilibra in MaTChM, taken from Zaveri (1997) and references therein.

No.	Dissociation Equilibrium	Dissociation Constant, M or M atm <sup>-1</sup>
D1.	$H_2O \iff H^+ + OH^-$	1.00E-14 exp(-6716FT)
D2.	$HO_2(1) \iff H^+ + O_2^-$	3.50E-5
D3.	$H_2O_2(1) \iff HO_2^- + H^+$	2.2E-12 exp(-3730FT)
D4.	$H_2CO_3 \iff HCO_3^- + H^+$	4.46E-7 exp(-1000FT)
D5.	$HCO_3^- \iff CO_3^{2-} + H^+$	4.48E-11 exp(-1760FT)
D6.	$HNO_3(1) \iff NO_3^- + H^+$	15.4
D7.	$HNO_2(1) \iff NO2^- + H^+$	5.1E-4 exp(-1260FT)
D8.	$NH_4OH \iff NH_4^+ + OH^-$	1.75E-5 exp(-450FT)
D9.	$HCHO(l) \iff CH_2(OH)_2(l)$	1.82E+3 exp(4020FT)
D10.	$H_2SO_3 \iff HSO_3^- + H^+$	1.23E-2 exp(1960FT)
D11.	$HSO_3^- \iff SO_3^{2-} + H^+$	6.61E-8 exp(1500FT)
D12.	$H_2SO_4(1) \iff HSO_4^- + H^+$	1.00E+3
D13.	$HSO_4^- \iff SO_4^{2-} + H^+$	1.02E-2 exp(2720FT)
D14.	$HCl(l) \iff H^+ + Cl^-$	1.74E+6 exp(6900FT)
D15.	$Cl_2^- \iff Cl + Cl^-$	5.26E-6
D16.	$HCOOH(I) \iff HCOO^- + H^+$	1.78E-4 exp(-20FT)
D17	$CH_3COOH(1) \iff CH_3COO^- + H^+$	1.0E-4
D18	CH <sub>3</sub> SO <sub>3</sub> H ⇔ CH <sub>3</sub> SO <sub>3</sub> <sup>-</sup> + H <sup>+</sup>	6.31E-2

Note: FT = (1/T - 1/298), T in Kelvin, Read 5.1E+6 exp(8700FT) as 5.1 x  $10^6$  e<sup>(8700(1//T-1/298))</sup>.